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(54) Title: **EXPRESSION PROFILING OF TUMOURS**

(57) Abstract: The present invention relates to methods of profiling tumours and characterisation of the tissue types associated with the tumours. A gene expression profile is obtained from the tissue sample, the genes ranked in order of their relative expression levels and the tissue type identified by comparing the gene ranking obtained with a database of relative gene expression level rankings of different tissue types. This gives a means to identify primary tumours and to determine the identity of a tumour of unknown primary. The invention also provides a method of treatment of a tumour by diagnosis of primary tumours identified by the methods described.

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Expression Profiling Of Tumors

The present invention relates to methods of profiling tumours and characterisation of the tissue types associated with the tumour. The present
5 invention also relates to a method of analysing gene expression data. Also provided is a means to identify primary tumours and to further determine the identity of a tumour of unknown primary. The invention also provides a method of treatment of a tumour by diagnosis of primary tumours identified by the methods described.

10

BACKGROUND

Advances in the treatment of cancer have resulted in significant improvements in median survival times for patients with many forms of the disease. These improvements have been the result of tailoring treatments to specific types of
15 tumours based on tissue specific molecular targets, for example hormone treatments for ovarian and breast cancers. However, if a tumour is misdiagnosed inappropriate treatment may delay recovery or have no effect on the disease. Therefore, there remains a need to correctly and reliably identify the source tissue of a tumour.

20

Despite the enormous amount of information regarding cancer and its diagnosis, there remains a significant proportion of new cancer cases that present with atypical symptoms. It has become apparent that the site of a tumour might belie its true origin. Metastatic tumours especially are in this class
25 because the primary tumour may be small or undetectable, consequently a large metastasis may be misdiagnosed as the primary tumour. Carcinomas of unknown origin account for between 3 and 5% of carcinomas. For example, a so-called Krukenberg tumour is a metastatic secondary carcinoma in the ovary and represents 6% of ovarian tumours. The primary tumour is usually a
30 mucinous carcinoma of the stomach. The definition is also broadly applied to tumours of the breast, pancreas and bowel metastatic to the ovary. Overall, approximately 20% of cancers in the ovary are thought to be of non-ovarian

origin. Figure 1 shows the most common sites of the primary in carcinoma of unknown primary in the cases where a primary is identified.

Current diagnostic techniques to identify the primary tumour in a patient with multiple disseminated metastases include morphological assessment, molecular pathological analyses including immunohistochemical staining, imaging techniques (CT, PET, mammography), and endoscopic techniques (gastroscopy, bronchoscopy, colonoscopy). While representing a small fraction of all patients, carcinoma of unknown primary accounts for the fourth most common cause of cancer death, mostly as the prognosis for these patients is bleak, with median survival eleven months. Carcinoma of unknown primary presents clinicians with a dilemma, namely how far to take investigation given the survival of patients with carcinoma of unknown primary is so poor. There is considerable debate concerning the value of detailed investigation to determine site of origin. Oncologists have been reluctant to perform low-yield investigations because of the unacceptable cost-effectiveness ratio. The cost of these investigations is not only monetary, but also impacts quality of life for the patient, and morbidity arising from invasive diagnostic procedures. Whether patients benefit from a more definitive diagnosis is unclear, however, it is the case that treatment approaches can vary significantly depending on cellular origin. For example, the drug therapies used for metastatic adenocarcinoma of the lung are significantly different to those used for metastatic adenocarcinoma of the pancreas, which, in turn are different to the therapeutic approach to metastatic colorectal adenocarcinoma.

Accordingly, it is desirable to provide a method to identify the origin of a tumour or the primary site in a carcinoma of an unknown primary so that effective treatment can be administered.

High-throughput expression analysis has recently been employed to great effect in the sub-classification of many tumour types. Since cancer is a disease of aberrant gene regulation, our ability to use microarrays to profile gene expression on a massively parallel level has begun to unravel the molecular

mechanisms behind tumour initiation, progression and response to therapy. The power of large-scale genetic analysis lies in the fact that the expressions of thousands of genes are used to characterise the tumour, rather than just several markers. Many examples now exist where formerly homogenous groupings of tumours based on conventional histopathological techniques have been subdivided into groups based on molecular profiling. Not unsurprisingly, the wealth of gene expression data in several diseases has begun to support the hypothesis that morphologically indistinguishable tumours may be molecularly distinguishable. This has potentially widespread application in the clinical application of technologies aimed at refining diagnosis and prognostication in cancer.

However, with the complex data derived from expression analysis, it is difficult to discern a meaningful result to fully diagnose and identify the primary tumour.

Accordingly, it is an aspect of the invention to provide a method to identify a primary site of a tumour.

A further difficulty encountered by those trying to identify a tumour's origin occurs when a patient develops a new tumour following an earlier disease. Typically such earlier disease will have been treated and a sample of the diseased tissue may have been stored by standard techniques such as paraffin embedding. In order to determine whether the new disease is related to the earlier disease it may be necessary to analyse gene expression in that archived sample. Conventional methods of gene expression analysis require high quality nucleic acid to be isolated, which is not possible from, for example, paraffin embedded tissue.

Thus, it is a further aspect of the present invention to provide a method of identifying a primary tumour from an archived or preserved sample.

SUMMARY OF THE INVENTION

In one aspect of the present invention, there is provided a method of profiling a biological sample, said method including:

- obtaining a gene expression profile from the biological sample;
- 5 obtaining a gene expression database from one or more biological samples;
- identifying different patterns of gene expression between the biological samples;
- identifying genes that comprise the different patterns of gene expression;
- 10 and
- correlating the genes that comprise the different patterns of gene expression of the gene expression profile of the biological sample and the gene expression database to provide a profile of the biological sample.

15 In another aspect of the present invention, there is provided a method of analysing gene expression data to generate a gene expression profile or a gene expression database for use in diagnosing tumours. Preferably the method allows comparison of data obtained from different experiments.

20 In yet another aspect of the present invention, there is provided a gene expression database generated using a method described herein.

In a further aspect of the present invention, there is provided an
25 expression-based diagnostic evaluation of the tissue of origin of a tumour. Preferably the expression-based evaluation is based on comparing a gene expression profile of a tumour with a gene expression database representing one or more tumour or tissue types.

30 In another aspect of the present invention, there is provided a method of treatment of a patient having a tumour of unknown origin including the steps of:
identifying the tissue of origin of the tumour of unknown origin; and

treating the patient in a manner appropriate for treating a tumour originating from that tissue.

5 An alternative gene expression profiling platform to cDNA microarray analysis is proposed using a system of high throughput RT-PCR (real time PCR). Key cancer class specific markers, identified through microarray analysis, can be easily translated to the RT-PCR method, allowing utilization of more robust and reproducible platform that could be integrated into a conventional pathology laboratory. Additionally, through using the method of RankLevels it has been
10 shown that microarray and RT-PCR datasets can be used for building integrated SVM predictor algorithms. This allows the utilization of datasets from both platforms for training and building such predictors. The RankLevel method can also be applied to cross platform meta-analysis to use or mine pre-existing gene expression datasets.

15

BRIEF DESCRIPTION OF THE FIGURES

Figure 1 shows the most common sites of the primary in carcinoma of unknown primary.

20 **Figure 2** shows the results of unsupervised hierarchical clustering of gene expression data from 121 primary tumours from a diverse range of human tumours.

Figure 3 shows a subset of genes which describes differences between tumour
25 types.

Figure 4 shows a graph indicating the results from the ranking of genes in order to identify a subset with the highest predictive strength.

30 **Figure 5** shows a confusion matrix constructed to show predictor accuracy as determined using the proportions of correct classifications from a leave-one-out cross validation in conjunction with a k-nearest neighbours algorithm.

Figure 6 shows the validity of the predictor algorithm by using it to identify the origin of twelve samples of metastatic tumour of unknown primary.

Figure 7 shows hierarchical clustering of ovarian (blue) and colorectal (red) primary tumours with Krukenberg-like tumours (green). All Krukenberg tumours co-cluster with colorectal primary tumour.

Figure 8 shows that support vector machine analysis with twelve tumour types identifies a colorectal source for the five Krukenberg-like tumours shown in **Figure 7**. The Y-axis represents a confidence measure of the prediction.

Figure 9 shows a heat map alignment of data generated using cDNA microarray and RT-PCR.

Figure 10 shows a hierarchical cluster analysis of RT-PCR data.

Figure 11 shows the performance of RankLevels for in classification of microarray data. Experiments presents accuracy of LOO (leave-one-out) cross validation on a set of 133 cancer samples divided into 16 classes. For **Figure A**, full precision of pin-group normalised expressions was used, for **Figures B** and **C** used RankLevels with 3 and 5 levels, respectively.

Figure 12 demonstrates the effect of dataset size and complexity on distribution of predictions within the three confidence levels and their relative accuracies. The *Complete* dataset represents LOOCV results from the complete dataset ($n=229$). *Training/Test* represents LOOCV results from Training set only ($n=167$). *LSO* represents the accumulated results from iteratively leaving subtypes from training ($n=96$). *LCO* represents accumulated results from iteratively leaving site of origin classes from training ($n=229$).

DETAILED DESCRIPTION OF THE INVENTION

In one aspect of the present invention, there is provided a method of profiling a biological sample, said method including:

- obtaining a gene expression profile from the biological sample;
- 5 obtaining a gene expression database from one or more biological samples;
- identifying different patterns of gene expression between the biological samples;
- identifying genes that comprise the different patterns of gene expression;
- 10 and
- correlating the genes that comprise the different patterns of gene expression of the gene expression profile of the biological sample and the gene expression database to provide a profile of the biological sample.

- 15 Applicants have used molecular profiling techniques to characterise tumours and various tissues of biological samples based on their gene expression profile. The underlying principle of this work is that an individual cell type only expresses a subset of the total number of genes present in the genome. The fraction of genes expressed reflects and determines the biological state of the
- 20 cell and provides a molecular snapshot of the cellular phenotype.

- As used herein the term "gene expression profile" includes information on the expression levels of a plurality of genes within a biological sample. A biological sample within the scope of the present invention may be any biological sample
- 25 that includes cellular material from which DNA, RNA or protein may be isolated. The expression level of a gene may be determined by the amount of DNA, RNA or protein present in the sample which corresponds with the gene. The gene expression profile therefore, may include levels of DNA, RNA and/or protein correlated to specific genes within the biological sample.

- 30 Gene expression levels may be obtained in a variety of ways including, but not limited to analysing DNA levels, mRNA levels, analysing protein levels and determining transcription initiation rates. Preferably gene expression levels are

determined by analysis of mRNA levels. More preferably mRNA levels are determined by a hybridisation-based method or a PCR-based method.

5 A variety of different biological samples may be used to generate a gene expression profile. For example, the biological sample may be a tissue sample and the tissue may be normal or diseased. A diseased tissue sample may include a pre-cancerous tissue, a cancerous tissue, a tumour, a primary tumour, a metastatic tumour, or cells collected from a pleural effusion. A pre-cancerous tissue includes a tissue which may become cancerous. The biological sample
10 may include freshly collected tissue, frozen tissue or archived tissue. In the case of archived tissue the sample may be a paraffin-embedded sample.

A gene expression profile may be established by hybridising a labelled nucleic acid sample from a biological sample to a plurality of target nucleic acids, and
15 detecting to which of the plurality of target nucleic acids the labelled nucleic acid has bound, thereby determining which of the plurality of target nucleic acids are expressed in the biological sample and establishing a gene expression profile for the biological sample. An exemplary method of gene expression analysis by a hybridisation-based technology includes the use of a microarray. In this
20 example, mRNA from a sample may be labelled either directly or through the synthesis of labelled cDNA. The labelled nucleic acid may then be hybridised to the microarray and expression levels determined by detecting the amount of labelled nucleic acid bound at particular positions on the microarray.

25 Alternatively or additionally, a PCR-based method of gene expression analysis may be used. For example, a quantitative RT-PCR technique. In this method, RNA from a biological sample may be reverse transcribed to generate segments of cDNA which may then be amplified by gene-specific quantitative PCR. The rate of accumulation of specific PCR products can be correlated to
30 the abundance of the corresponding RNA species in the original sample and thereby provide an indication of gene expression levels. An RT-PCR method of gene expression analysis provides a robust method for obtaining expression data in a short time, compared with hybridisation-based techniques.

Both of the aforementioned techniques determine the expression of a gene by measuring the amount of mRNA corresponding to the gene.

- 5 Protein expression data may also be included in a gene expression profile since the level of a protein product generally represents the functional expression level of a gene. Protein expression levels may be determined by a hybridisation assay such as binding to an antibody or other ligand, or a functional assay where a specific protein function or activity may be measured directly.

10

Although less suited to high throughput or rapid analysis, transcription initiation rates may also provide an indication of gene expression levels. Such analyses require the use of a living sample in which nascent RNA transcripts are pulse labelled in vivo and analysed in a gene specific manner, generally involving
15 hybridisation to unlabelled target nucleic acid representing the gene of interest. The labelled RNA only represents genes being actively transcribed and gives an indication of the rate of transcription initiation of a gene.

20

Hence a gene expression profile provides information on the expression level of a plurality of genes within a biological sample. Preferably the biological sample is a tissue sample. More preferably the biological sample is a tumour sample. The tumour sample may be of known origin or of unknown origin.

25

In particular embodiments of the present invention a plurality of gene expression profiles may be used to generate a gene expression database.

30

As used herein the term "gene expression database" refers to the expression profiles for a given sample type or types. A plurality of gene expression profiles may be used to generate the gene expression database. The gene expression profiles are statistically analysed to identify gene expression levels that characterise particular sample types. The gene expression database may also be established for a given tissue type or plurality of tissue types, and thus, in particular embodiments of the present invention, may allow the identification of

the tissue from which a tumour was originally derived, by comparing the tumour's gene expression profile to the gene expression database.

Hence a gene expression database establishes a "fingerprint" of the expression profiles for a given sample type. Preferably the sample is a tissue sample.
5 More preferably the sample is a tumour sample. In particular embodiments of the present invention a gene expression database includes gene expression information for one or more sample types, including but not limited to any one or more of the following tumours: gastric, colorectal, pancreatic, breast and
10 ovarian.

Patterns of gene expression may be determined by statistical analysis of a gene expression profile or a gene expression database. Preferably the analysis employs an algorithm which utilises a number of informatic tools including k-
15 nearest neighbours and a support vector machine (SVM) approach. In analyzing gene expression data, the first stage is to reduce the number of genes analysed to an optimal subset, capable of reliably describing differences between tumour types. This step is necessary as microarray-derived gene expression profiles may include data from the many thousands of genes
20 represented on the array. Preferably, an initial step of normalizing the data is employed. Depending on the method by which the expression data is obtained, the normalization procedure may be accompanied by a Ranking System, described below. Generally, with microarray data, the number of data points is large and normalization is needed to reduce the numbers and exclude noise
25 and aberrant data to a manageable level. However, when using RT_PCR to generate the gene expression data, the number of data points is much less and hence more manageable. Therefore, these datapoints may undergo a Ranking process at this stage as described below.

30 The optimal number and selection of genes for classification of tumours and biological samples from a range of primary origins is determined by using an iterative signal to noise ratio algorithm. This method ranks genes according to the difference of their mean expression values for each class of tumour, divided

by the sum of the standard deviations, ie. $(m_1 - m_2)/(s_1 + s_2)$. This effectively identifies those genes that have a consistently different expression measurement within a given class of tumours, relative to the values of that gene across all other tumour types present. This method may also be employed
5 when RT-PCR is used to validate the gene expression profiles of the samples. For, instance, a microarray may be used to initially test a number of genes from which a reduced set of expressing genes indicative of the sample may then be applied to an assay such as RT-PCR which requires less gene sets (but more specific genes) and generates fewer data points.

10

To select and test subsets of genes, a leave-one-out (LOO) cross validation in conjunction with the k-nearest neighbors algorithm can be used. Briefly, this algorithm seeks to classify an unknown sample by comparing it to samples of known class by using a distance metric. The class of the closest 'k' samples is
15 assigned to the sample being tested. LOO involves permutations of the dataset whereby each sample is held out separately and a class assigned to it by using the remaining samples. This is repeated until each sample has been left out of the training set once and been assigned to a class. The proportion of correct classifications is used as a measure of predictor accuracy. By plotting the actual
20 tumour classes on one axis and the predicted classes on the other, a histogram-type view of the overall success or failure of the classification approach can be achieved. This representation (see for example Figure 5) also allows identification of any particular classes with more incorrect predictions relative to other tumour types. The average prediction accuracy in LOO
25 analysis in one particular training set is approximately 97%.

The applicants have generated a training set of over 120 primary tumours from a diverse range of human tumours, representing the major tumour types accounting for carcinoma of unknown primary (see Table 1). Unsupervised
30 hierarchical clustering of gene expression data from these tumours results in a near perfect segregation of different tumour types. Figure 2 shows the results of such a cluster, with approximately 500 genes selected on the basis of at least three samples with an expression ratio greater than or equal to 2.7.

Table 1. Summary of tumour samples used in training set.

Key: test: samples processed on MFC, trai: samples processed by microarray not by MFC
 trai2: samples not used for MFC

Patient ID	CancerType	DataClass	Comment
P00030	breast	test	Primary ER Positive
P00640	breast	test	Metastasis ER Positive
P00734	breast	test	Primary
P00743	breast	test	Primary ER Positive
P01026	breast	test	Primary ER Negative
P01212	breast	test	Primary ER Positive
P01374	breast	test	Primary ER Positive
P01398	breast	test	Primary ER Positive
P01696	breast	test	Primary ER Positive
P02274	breast	test	Primary ER Positive
P02288	breast	test	Primary ER Positive
P00541	colorectal	test	Primary
P00617	colorectal	test	Primary
P01740	colorectal	test	Primary
P01757	colorectal	test	Primary
P01840	colorectal	test	Primary
P02225	colorectal	test	Primary
P02553	colorectal	test	Primary
P02740	colorectal	test	Primary
P00448	Gastric	test	Primary Intestinal
P00514	Gastric	test	Primary Intestinal
P00553	Gastric	test	Primary Diffuse
P00559	Gastric	test	Primary Intestinal
P00628	Gastric	test	Primary Intestinal
P00661	Gastric	test	Primary Signet ring
P02173	Gastric	test	Primary Diffuse
P02176	Gastric	test	Primary Diffuse
P02318	Gastric	test	Primary Diffuse
P00195	ovarian	test	Primary serous
P00446	ovarian	test	Primary serous
P00633	ovarian	test	Primary serous
P00756	ovarian	test	Primary serous
P00772	ovarian	test	Primary serous
P01164	ovarian	test	Metastasis serous
P01246	ovarian	test	Primary serous
P01428	ovarian	test	Primary serous
P01436	ovarian	test	Primary serous
P02244	pancreas	test	Primary
P02245	pancreas	test	Primary
P02246	pancreas	test	Primary
P02248	pancreas	test	Primary
P02249	pancreas	test	Primary
P02250	pancreas	test	Primary
P03056	pancreas	test	Primary
P00006	breast	trai	Primary ER Positive
P00009	breast	trai	Primary ER Positive
P00066	breast	trai	Primary ER Negative

P00442	breast	tra	Primary ER Negative
P00467	breast	tra	Primary ER Positive
P00469	breast	tra	Primary ER Negative
P00478	breast	tra	Primary ER Positive
P00504	breast	tra	Primary ER Positive
P00546	breast	tra	Primary ER Negative
P00572	breast	tra	Primary ER Positive
P00621	breast	tra	Primary ER Positive
P00746	breast	tra	Primary ER Negative
P00776	breast	tra	Primary ER Negative
P00786	breast	tra	Primary ER Positive
P00905	breast	tra	Primary ER Negative
P00993	breast	tra	Primary ER Negative
P01289	breast	tra	Metastasis ER Positive
P01292	breast	tra	Primary ER Negative
P01579	breast	tra	Metastasis ER Positive
P01843	breast	tra	Primary ER Negative
P01944	breast	tra	Metastasis ER Negative
P00002	colorectal	tra	Primary
P00049	colorectal	tra	Primary
P00578	colorectal	tra	Primary
P00587	colorectal	tra	Primary
P00721	colorectal	tra	Metastasis
P00759	colorectal	tra	Metastasis
P00896	colorectal	tra	Metastasis
P00961	colorectal	tra	Primary
P00967	colorectal	tra	Metastasis Sigmoid
P00974	colorectal	tra	Primary
P01016	colorectal	tra	Metastasis
P01060	colorectal	tra	Metastasis
P01838	colorectal	tra	Primary
P01844	colorectal	tra	Metastasis
P01905	colorectal	tra	Primary
P00035	Gastric	tra	Primary Diffuse
P00048	Gastric	tra	Metastasis Signet ring
P00051	Gastric	tra	Primary Signet ring
P00433	Gastric	tra	Primary Diffuse
P00483	Gastric	tra	Primary Diffuse
P00503	Gastric	tra	Metastasis Diffuse
P00536	Gastric	tra	Primary Diffuse
P00551	Gastric	tra	Primary Diffuse
P00109	ovarian	tra	Primary endometriod
P00130	ovarian	tra	Primary serous
P00151	ovarian	tra	Primary endometriod
P00155	ovarian	tra	Primary serous
P00160	ovarian	tra	Primary serous
P00164	ovarian	tra	Primary endometriod
P00165	ovarian	tra	Primary serous
P00169	ovarian	tra	Primary mucinous
P00188	ovarian	tra	Primary endometriod
P00488	ovarian	tra	Primary mucinous
P00496	ovarian	tra	Metastasis
P00505	ovarian	tra	Primary endometriod

P00506	ovarian	tra	Primary endometriod
P00511	ovarian	tra	Primary serous
P00627	ovarian	tra	Primary mucinous
P00706	ovarian	tra	Metastasis serous
P00718	ovarian	tra	Primary mucinous
P00784	ovarian	tra	Primary mucinous
P00807	ovarian	tra	Primary mucinous
P00809	ovarian	tra	Primary serous
P00933	ovarian	tra	Primary serous
P00935	ovarian	tra	Primary mucinous
P01348	ovarian	tra	Primary serous
P01563	ovarian	tra	Primary serous
RBH 91	ovarian	tra	Primary endometriod RBH 91.033
RBH 92	ovarian	tra	Primary mucinous RBH 92.011
RBH 93	ovarian	tra	Primary endometriod RBH 93.118
RBH 93	ovarian	tra	Primary endometriod RBH 93.061
RBH 93	ovarian	tra	Primary mucinous RBH 93.002
RBH 93	ovarian	tra	Primary mucinous RBH 93.085
RBH 94	ovarian	tra	Primary endometriod RBH 94.037
RBH 94	ovarian	tra	Primary endometriod RBH 94.120
RBH 94	ovarian	tra	Primary endometriod RBH 94.020
RBH 94	ovarian	tra	Primary mucinous RBH 94.030
RBH 94	ovarian	tra	Primary mucinous RBH 94.072
RBH 94	ovarian	tra	Primary mucinous RBH 94.080
WM 090	ovarian	tra	Primary malignant mucinous
WM 223	ovarian	tra	Primary mucinous
WM 438	ovarian	tra	Primary mucinous
WM 439	ovarian	tra	Primary mucinous
WM 454	ovarian	tra	Primary malignant mucinous
P02078	pancreas	tra	Primary
P02247	pancreas	tra	Primary
P00815	Lung	tra2	Primary sc
P00817	Lung	tra2	Primary sc
P00925	Lung	tra2	Primary sc
P01323	Lung	tra2	Primary sc
P01400	Lung	tra2	Primary sc
P01759	Lung	tra2	Primary adenocarcinoma
P01770	Lung	tra2	Primary adenocarcinoma
P01907	Lung	tra2	Primary sc
P01909	Lung	tra2	Primary adenocarcinoma
P02021	Lung	tra2	Primary sc
P02023	Lung	tra2	Primary adenocarcinoma
P02024	Lung	tra2	Primary large cell
P02025	Lung	tra2	Primary adenocarcinoma
P02026	Lung	tra2	Primary adenocarcinoma
P02028	Lung	tra2	Primary large cell
P02029	Lung	tra2	Primary sc
P02030	Lung	tra2	Primary large cell
P02031	Lung	tra2	Primary sc
P02032	Lung	tra2	Primary adenocarcinoma
P02033	Lung	tra2	Primary adenocarcinoma
P02034	Lung	tra2	Primary large cell
P02035	Lung	tra2	Primary adenocarcinoma

P02037	Lung	tra2	Primary adenocarcinoma
P02038	Lung	tra2	Primary adenocarcinoma
P02039	Lung	tra2	Primary sc
P02040	Lung	tra2	Primary large cell
P02041	Lung	tra2	Primary large cell
P02042	Lung	tra2	Primary large cell
P02043	Lung	tra2	Primary sc
P02044	Lung	tra2	Primary sc
P02045	Lung	tra2	Primary large cell
P02090	Lung	tra2	Primary large cell
P00508	melanoma	tra2	Metastasis
P00576	melanoma	tra2	Metastasis
P00761	melanoma	tra2	Metastasis
P00825	melanoma	tra2	Metastasis
P00833	melanoma	tra2	Metastasis
P00923	melanoma	tra2	Metastasis
P00977	melanoma	tra2	Metastasis
P00979	melanoma	tra2	Metastasis
P01537	melanoma	tra2	Metastasis
P01861	melanoma	tra2	Metastasis
P01726	mesothelioma	tra2	Primary
P01728	mesothelioma	tra2	Primary
P01729	mesothelioma	tra2	Primary
P01730	mesothelioma	tra2	Primary
P01731	mesothelioma	tra2	Primary
P01733	mesothelioma	tra2	Primary
P00050	Oesophageal	tra2	Primary Mixed
P00450	Oesophageal	tra2	Primary Diffuse
P00032	prostate	tra2	Primary
P00880	prostate	tra2	Primary
P00890	prostate	tra2	Primary
P00954	prostate	tra2	Primary
P01109	prostate	tra2	Primary
P01421	prostate	tra2	Primary
P01653	prostate	tra2	Primary
P01813	prostate	tra2	Primary
P00916	renal	tra2	Metastasis
P00998	renal	tra2	Primary
P01020	renal	tra2	Primary
P01038	renal	tra2	Primary
P01043	renal	tra2	Primary
P01048	renal	tra2	Primary Clear cell
P01098	renal	tra2	Primary
P01218	renal	tra2	Primary
P01270	renal	tra2	Primary Clear cell
P01278	renal	tra2	Primary
P01574	renal	tra2	Metastasis
P01817	renal	tra2	Primary
P01908	renal	tra2	Metastasis
P01093	SCCo	tra2	Primary Larynx, NOS
P01158	SCCo	tra2	Primary Tongue, NOS
P01308	SCCo	tra2	Primary Tongue, NOS
P01343	SCCo	tra2	Primary Pharynx, NOS

P01394	SCCoother	tra12	Primary Pyriform sinus
P01472	SCCoother	tra12	Primary Larynx, NOS
P01749	SCCoother	tra12	Primary Larynx, NOS
P01273	SCCoother Skin	tra12	Primary Skin of lip, NOS
P01341	SCCoother Skin	tra12	Unknown Unknown primary site
P01371	SCCoother Skin	tra12	Primary Skin, NOS
P01402	SCCoother Skin	tra12	Primary Skin of scalp and neck
P01633	SCCoother Skin	tra12	Primary Skin of other and unspecified parts of face
P01660	SCCoother Skin	tra12	Primary Skin, NOS
P01766	SCCoother Skin	tra12	Primary Skin of other and unspecified parts of face
P01832	SCCoother Skin	tra12	Unknown Unknown primary site
P00876	testicular	tra12	Primary
P01124	testicular	tra12	Primary
P02345	testicular	tra12	Primary
P00635	uterine	tra12	Primary endometriod
P00724	uterine	tra12	Primary endometriod
P00741	uterine	tra12	Primary endometriod
P00742	uterine	tra12	Primary endometriod
P00848	uterine	tra12	Primary endometriod
P00909	uterine	tra12	Primary endometriod
P00940	uterine	tra12	Primary endometriod
P00943	uterine	tra12	Primary endometriod
P01872	uterine	tra12	Primary endometriod

In a preferred embodiment of the present invention there is provided a set of approximately 90 genes (see Table 2 below), many of which may be used for discriminating between a plurality of sample types, including but not limited to any one or more of the following tumours: gastric, colorectal, pancreatic, breast and ovarian.

Table 2. A set of genes useful in discriminating gastric, colorectal, pancreatic, breast and ovarian tumours.

Genbank	RefSeq	Name	Symbol	Class
AA291749	NM_000125	estrogen receptor 1	ESR1	brea
AA479494	NM_020423	ezrin-binding partner PACE-1	PACE-1	brea
AA479888	NM_004703	rabaptin-5	RAB5EP	brea
AA482035	NM_014804	KIAA0753 gene product	KIAA0753	brea
AA489647	NM_004354	cyclin G2	CCNG2	brea
AI362703	NM_007255	xylosylprotein beta 1,4-galactosyltransferase, polypeptide 7 (galactosyltransferase I)	B4GALT7	brea
AI635773	NM_025202	likely ortholog of neuronally expressed calcium binding protein	FLJ13612	brea
AI669721	NM_014112	trichorhinophalangeal syndrome I	TRPS1	brea
AI972286	NM_002652	prolactin-induced protein	PIP	brea
H10045	NM_006113	vav 3 oncogene	VAV3	brea
H29315	NM_012319	LIV-1 protein, estrogen regulated	LIV-1	brea
H72875	NM_002051	GATA binding protein 3	GATA3	brea
N23299	NM_014674	ER degradation enhancing alpha mannosidase-like	EDEM	brea
N49284	NM_005375	v-myb myeloblastosis viral oncogene homolog (avian)	MYB	brea
R06567	NM_003629	phosphoinositide-3-kinase, regulatory subunit, polypeptide 3 (p55, gamma)	PIK3R3	brea
R63647	NM_000949	prolactin receptor	PRLR	brea
H95792	NM_001609	acyl-Coenzyme A dehydrogenase, short/branched chain	ACADSB	brea
AA088420	NM_015869	peroxisome proliferative activated receptor, gamma	PPARG	colo
AA099136	NM_002296	lamin B receptor	LBR	colo
AA130579	NM_006149	lectin, galactoside-binding, soluble, 4 (galectin 4)	LGALS4	colo
AA130584	NM_004363	carcinoembryonic antigen-related cell adhesion molecule 5	CEACAM5	colo
AA262074	NM_147130	natural cytotoxicity triggering receptor 3	NCR3	colo
AA279081	NM_015250	coiled-coil protein BICD2	BICD2	colo
AA284184	NM_018438	F-box only protein 6	FBXO6	colo
AA406571	NM_001712	carcinoembryonic antigen-related cell adhesion molecule 1 (biliary glycoprotein)	CEACAM1	colo
AA465495	NM_016234	fatty-acid-Coenzyme A ligase, long-chain 5	FACL5	colo
AA699679	NM_003889	nuclear receptor subfamily 1, group I, member 2	NR1I2	colo
AA975612	NM_012396	pleckstrin homology-like domain, family A, member 3	PHLDA3	colo
AI433336	NM_007127	villin 1	VIL1	colo
AI681730	NM_007052	NADPH oxidase 1	NOX1	colo

AW009320	NM_001804	caudal type homeo box transcription factor 1	CDX1	colo
N74131	NM_003226	trefoil factor 3 (intestinal)	TFF3	colo
W72792	NM_004442	EphB2	EPHB2	colo
AA490044	NM_006933	solute carrier family 5 (inositol transporters), member 3	SLC5A3	gast
AA664101	NM_000689	aldehyde dehydrogenase 1 family, member A1	ALDH1A1	gast
AA702350	NM_015570	autism susceptibility candidate 2	AUTS2	gast
AA702640	NM_000790	dopa decarboxylase (aromatic L-amino acid decarboxylase)	DDC	gast
AA845156	NM_003122	serine protease inhibitor, Kazal type 1	SPINK1	gast
AI090702	NM_014970	kinesin-associated protein 3	KIFAP3	gast
AI333599	NM_019617	18 kDa antrum mucosa protein	AMP18	gast
AW009769	NM_003225	trefoil factor 1 (breast cancer, estrogen-inducible sequence expressed in)	TFF1	gast
AW029441	NM_002630	progastricsin (pepsinogen C)	PGC	gast
AW058221	NM_004190	lipase, gastric	LIPF	gast
H23187	NM_000067	carbonic anhydrase II	CA2	gast
H94487	NM_001910	cathepsin E	CTSE	gast
N63943	NM_000239	lysozyme (renal amyloidosis)	LYZ	gast
R32848	NM_005980	S100 calcium binding protein P	S100P	gast
R39069	NM_003558	phosphatidylinositol-4-phosphate 5-kinase, type I, beta	PIP5K1B	gast
T60861	NM_017846	tRNA selenocysteine associated protein	SECP43	gast
AA405767	NM_013952	paired box gene 8	PAX8	ovar
AA419229	NM_144586	hypothetical protein MGC29643	MGC29643	ovar
AA453742	NM_004172	solute carrier family 1 (glial high affinity glutamate transporter), member 3	SLC1A3	ovar
AA459363	NM_017495	RNA-binding region (RNP1, RRM) containing 1	RNPC1	ovar
AA621342	NM_015415	DKFZP564B167 protein	DKFZP564B167	ovar
AA683520	NM_003064	secretory leukocyte protease inhibitor (antileukoprotease)	SLPI	ovar
AI139437	NM_005046	kallikrein 7 (chymotryptic, stratum corneum)	KLK7	ovar
AI963941	NM_144505	kallikrein 8 (neuropsin/ovasin)	KLK8	ovar
N52450	NM_033624	F-box only protein 21	FBXO21	ovar
R24530	NM_016730	folate receptor 1 (adult)	FOLR1	ovar
AA122287	NM_005512	glycoprotein A repetitions predominant	GARP	panc
AA450265	NM_002592	proliferating cell nuclear antigen	PCNA	panc
AA454651	NM_020831	megakaryoblastic leukemia (translocation) 1	MKL1	panc
AA670378	NM_014504	putative Rab5 GDP/GTP exchange factor homologue	RABEX5	panc
AA670429	NM_003020	secretory granule, neuroendocrine protein 1 (7B2 protein)	SGNE1	panc

AA844864	NM_006507	regenerating islet-derived 1 beta (pancreatic stone protein, pancreatic thread protein)	REG1B	panc
AA845178	NM_001868	carboxypeptidase A1 (pancreatic)	CPA1	panc
AA894687	NM_004515	interleukin enhancer binding factor 2, 45kDa	ILF2	panc
AI651194	NM_015089	p53-associated parkin-like cytoplasmic protein	PARC	panc
AI669320	NM_006418	differentially expressed in hematopoietic lineages	GW112	panc
AI685081	NM_000207	insulin	INS	panc
AI829222	NM_000371	transferrin (prealbumin, amyloidosis type I)	TTR	panc
T54662	NM_001832	collipase, pancreatic	CLPS	panc
W45219	NM_006229	pancreatic lipase-related protein 1	PNLIPRP1	panc
W72322	NM_001419	ELAV (embryonic lethal, abnormal vision, Drosophila)-like 1 (Hu antigen R)	ELAVL1	panc
AA400464	NM_000346	SRY (sex determining region Y)-box 9 (campomelic dysplasia, autosomal sex-reversal)	SOX9	oth
AA402040	NM_014428	tight junction protein 3 (zona occludens 3)	TJP3	oth
AA430665	NM_001305	claudin 4	CLDN4	oth
AA443558	NM_032420	protocadherin 1 (cadherin-like 1)	PCDH1	oth
AA477165	NM_002906	radixin	RDX	oth
AA676466	NM_054012	argininosuccinate synthetase	ASS	oth
AA872020	NM_002773	protease, serine, 8 (protease)	PRSS8	oth
AA972350	NM_000542	surfactant, pulmonary-associated protein B	SFTPB	oth
AI002217	NM_003019	surfactant, pulmonary-associated protein D	SFTPD	oth
N58558	NM_006215	serine (or cysteine) proteinase inhibitor, clade A (alpha-1 antiproteinase, antitrypsin), member 4	SERPINA4	oth
N68998	NM_014382	ATPase, Ca++ transporting, type 2C, member 1	ATP2C1	oth
R99562	NM_004497	forkhead box A3	FOXA3	oth
AA001444	NM_002398	Meis1, myeloid ecotropic viral integration site 1 homolog (mouse)	MEIS1	ovar
AA130187	NM_024426	Wilms tumour 1	WT1	ovar
AA142980	NM_015470	gamma-SNAP-associated factor 1	GAF1	ovar
H89996	NM_006565	CCCTC-binding factor (zinc finger protein)	CTCF	control
AA430524	NM_004930	capping protein (actin filament) muscle Z-line, beta	CAPZB	control
AA078976	NM_004786	thioredoxin-like, 32kDa	TXNL	control
AA421230	NM_012433	splicing factor 3b, subunit 1, 155kDa	SF3B1	control
AA456028	NM_004582	Rab geranyltransferase, beta subunit	RABGGTB	control
AA419281	NM_002046	glyceraldehyde-3-phosphate dehydrogenase	GAPD	control

An alternative or complementary method for analysis of a gene expression database uses analyses similar to those described above to identify a subset of informative genes which may be used to discriminate between various sample types. For example a subset of approximately 90 genes may be used to discriminate between five classes of tumours: gastric, colorectal, pancreatic, breast and ovarian. Expression levels of each of those genes may then be ranked within each sample type thus resulting in an ordered list of genes that may be used to discriminate between different samples based on the relative expression levels of specific genes. This is known as Ranking, as herein described. This method has particular application and utility as it provides a method by which a sample may be identified without reference to a database. In a preferred embodiment a sample may be analyzed for expression levels of a specific set of genes, the relative expression levels of those genes may then be determined and ranked, then compared to a listing generated from different samples on the same set of genes, thereby providing a simple method of identifying the sample. This ranking procedure allows for meta-analysis which provides for cross-platform comparisons of gene expression profiles and databases.

In another aspect of the present invention, there is provided a method of analysing gene expression data to generate a gene expression profile or a gene expression database for use in diagnosing tumours. Preferably the method uses normalising gene expression data which allows comparison of data obtained from different experiments.

The methods described above relating to the generation and analysis of a gene expression profile or a gene expression database will now be described in more detail in this specific example. However, this application is not limited to this description and should not limit the generality of this invention.

As the present invention may use data generated from a variety of gene expression analysis methods including, but not limited to, microarray analysis and RT-PCR, a statistical method is required which facilitates amalgamation of

these data into a form which allows comparison of these different data. Applicants have also developed a Ranking System which is a surprisingly straightforward and robust approach to gene expression analysis.

5 Current microarray based measurements of gene expression are very noisy. This applies in particular to spotted array technologies used for development of this invention. The current dogma is that the raw measurement values have to be accordingly normalised, then various machine learning techniques should/could be applied to the normalised expression levels. A particular aim of
10 the normalisation is to combat the noise and some innate biases of the technology, such as the non-linear dependence between intensity and level of hybridisation of the Cy3 and Cy5 channels, wherein, Cy3 and Cy5 are fluorescent dyes used to label probes for the detection of nucleic acid hybridisation to microarrays. A number of sophisticated statistical normalisation
15 techniques were custom designed to suit various microarray platforms and results of their experimental evaluations can be found in the literature. These include the intensity dependent loess pin group normalisation for spotted arrays of Yang *et al* (2002, *Nucleic Acids Res* 30(4): e15), the SNOMAD algorithm of Colantuoni *et al* (2002, *Biotechniques* 32(6): 1316-20) for spatial normalisation
20 of spotted arrays, standardisation of gene expression values for Affymetrix array data to zero mean and unit standard deviation and universally applied log transformation alleviating routinely observed large dynamic ranges of expression values.

25 The present invention introduces, in particular, a novel normalisation technique based on ranking. Applicants propose to rank all genes according to their expression levels, then allocate to each gene a rough level of its rank (*RankLevel*). The *RankLevels* are then used for statistical analysis and predictive modelling instead of using normalised expression levels.

30

Effectively, raw expression data is obtained which provides a gene expression profile. This raw expression data may be obtained by microarray or RT-PCR analysis or any means that provides gene expression data. This data is

preferably normalized and reduced to a manageable level before processing through a k-nearest neighbours or SVM procedure or any learning algorithm process which is trained from the the data in a gene expression database. The ranking system, described herein, ranks the expression levels of the various data points within a sample.

Each data point represents an expression of a gene and is measured by the relative abundance of mRNA species in the sample compared to expression of that gene in a reference sample or median expression across many samples or genes. The intensity is assigned an intensity level which is determined relative to a reference point such as the background. Hence an intensity ratio or expression ratio may be obtained which represents the data point. This intensity or expression ratio is then ranked along with other data points within the sample ranging from the lowest intensity to the highest intensity. Each data point is then allocated a rank number. It is this rank number which is used to determine the rank level.

More specifically, if the expression ratio of a gene A is say 1.883 and gene B is 10.34, these values may be ranked as 5405 and 7283 among all 8378 genes of the array ordered from the lowest to the highest gene expression. Therefore, they are the 5405th and the 7283rd intense genes of the 8378 genes analysed. A random number of rank levels may be assigned. Preferably, these are ranked from 1 to 10. However, any number may be assigned. Useful ranks are between 2 to 10, most preferably 4 to 10, more preferably, 5 to 10. Assuming 5 RankLevels are used, then these genes are allocated RankLevels according to the following formula:

$$\begin{aligned} \text{RankLevel}(A) &= \text{ceil}(5 * 5405 / 8378) = \text{ceil}(3.22) = 4, \\ \text{RankLevel}(B) &= \text{ceil}(5 * 7283 / 8378) = \text{ceil}(4.34) = 5, \end{aligned}$$

where $\text{ceil}(x)$ denotes smallest integer $\geq x$. Thus, for building a predictive model based on RankLevels the values 4 and 5 can be used instead of the original values 1.883 and 10.34 for the expression of gene A and B,

respectively. Surprisingly, in spite of its crudeness, the RankLevel value allows development of very accurate predictive models (Figure 11). The intuitive explanation is that for successful predictive modelling the consistency of the features used to represent measurement is paramount. Thus although

5 RankLevels may lose accuracy of expression level, they gain in stability, since crude RankLevels are more likely to be unchanged or not changed significantly between various experiments. The RankLevels naturally eliminate the issue of huge dynamic range of expression values and global variations between average expression levels of different microarray slides. Hence a general

10 formula for determining a RankLevel is as follows:

$$\begin{aligned} \text{RankLevel} &= \text{ceil} (\text{number of rank levels} \times \\ &\quad \text{rank/number of genes assayed}) \\ &= \text{ceil} (x) \end{aligned}$$

wherein $\text{ceil} (x) = \text{smallest integer} \geq x$

15

Another important property of RankLevels is that models built on them can be easily transferred across various technologies for measuring gene expression levels, as long as the monotonicity of measurement across the technological platforms is roughly preserved. In various experiments applicants have very

20 successfully transferred predictive models developed for spotted microarrays to RT-PCR and vice versa (Example 7 and Figure 9). The same transfer can be done between other platforms, for instance, the spotted arrays and Affymetrix arrays. Note also that RankLevels are readily applicable to a mixture of arrays with different total number of genes that paves an avenue to practical statistical

25 analysis and modelling across large amounts of data from a variety of studies developed by different laboratories using various technologies.

RankLevel based models (using small number of rank levels, say 2-5) are also very amenable to human comprehension and rationalisation that can be readily

30 carried across range of technological platforms. In this context RankLevel normalization is especially attractive proposition, for emerging applications of microarray and RT-PCR technology and for other high throughput genetic experiments and their applications.

High-throughput expression analysis can be employed to great effect in the sub-classification of many tumour types. The wealth of gene expression data in several diseases has begun to support the hypothesis that morphologically
5 indistinguishable tumours may be molecularly distinguishable. This has potentially widespread application in the clinical application of technologies aimed at refining diagnosis and prognostication in cancer.

10 In yet another aspect of the present invention, there is provided a gene expression database generated using a method described herein. Preferably, the gene expression database includes a subset of genes selected to demonstrate differences in expression between sample types, and arranged according to their RankLevel as described above.

15 In a preferred embodiment the present invention provides a gene expression database which includes gene expression data from a variety of tumour types. Preferably the tumour types includes at least one of the following: gastric, colorectal, pancreatic, breast and ovarian.

20 In a further aspect of the present invention, there is provided an expression-based diagnostic evaluation of the tissue of origin of a tumour. Preferably the expression-based evaluation is based on comparing a gene expression profile of a tumour with a gene expression database representing one or more tumour or tissue types. More preferably, the comparison is based
25 on comparing RankLevels between the gene expression profile and the gene expression database.

Being able to provide disease appropriate treatment is essential in order to provide the best level of care for a patient. Given that different tumour types
30 respond differently to different treatment regimens, it is therefore beneficial to be able to correctly diagnose a patient's tumour. At present in medicine, the ability to classify tumours is based upon the use of a limited number of markers, which are often thought to be "tumour specific" in expression but in practice may

produce equivocal results regarding the tissue of origin of a tumour sample. For instance, although the estrogen receptor is employed as a diagnostic marker for breast cancer, the molecule is expressed in only a small percentage of clinically identifiable breast cancer samples. To further complicate the analysis, the
5 estrogen receptor is also expressed in various other tumour types. Thus present diagnosis is based on a limited set of imperfect predictors.

As stated above, the fraction of genes expressed in a cell reflects and determines the biological state of that cell and provides a molecular snapshot of
10 the cellular phenotype. Despite being propagated for many years in vitro, cell lines retain some level of lineage specific expression. This has the effect of allowing cell lines of similar origin to co-cluster following gene expression analyses. In addition, expression profiles of tumour cells in vivo or in vitro may group the cells according to their presumptive tissue of origin. Our ability to
15 rapidly profile the expression of many thousands of genes simultaneously, and use that information to diagnose the origin of a tumour has as yet not been reflected in modern diagnostics. The power of molecular profiling as an approach to diagnostic evaluation of tumours lies in the fact that instead of deriving information about a tumour from a handful of markers, the expression
20 of thousands of genes contributes to an overall picture of the tumour cells. The present invention confirms the diagnostic utility of such an approach, and foreshadows an expanding use of this technology. Preferably the expression-based evaluation uses expression data generated by the use of microarray technology to determine RNA expression levels in a sample.
25 Alternatively or additionally, the expression-based evaluation uses expression data generated by the use of quantitative RT-PCR technology to determine RNA expression levels in a sample.

The use of microarrays and quantitative RT-PCR generates a large amount of
30 data and requires considerable analysis to identify an optimal subset of genes, as discussed above. Once an optimal subset of genes has been identified, it is only necessary to investigate those genes in the optimal subset in order to perform identification according to the present invention.

In a particular embodiment of the present invention there is provided a method by which a tissue of origin or a tumour of origin may be assigned to a biological sample, the method including the steps of:

- 5 obtaining a gene expression profile of the biological sample; and
 comparing the gene expression profile to a gene expression database;
 wherein the gene expression database includes gene expression data
 relating to various tissue types or tumour types;
 wherein similarities and differences between the gene expression profile and
10 the gene expression database allow assignment of the tissue of origin or the
 tumour of origin to the biological sample.

In a preferred embodiment the biological sample is a tumour sample. More preferably the tumour sample is an unidentified adenocarcinoma. Preferably
15 the gene expression database includes gene expression data relating to any one or more of the following tumour types: gastric, colorectal, pancreatic, breast and ovarian.

Thus the present invention provides a method of diagnosing a patient's tumour
20 by comparing a gene expression profile of the patient's tumour with a gene expression database generated from known tumour types.

In a particular embodiment of the present invention the methods of the invention can be used to identify a tumour of unknown origin.

25 In a specific, but non-limiting example, the present application illustrates the process in the identification of tumours found in the ovary, but suspected to be extra-ovarian in origin. Approximately 10-20% of patients presenting with ovarian malignancies have tumours suspected to be of extra-ovarian origin,
30 rather than primary ovarian cancers. Tumours that metastasise from the stomach to the ovary and present as primary ovarian cancer are typically referred to as Krukenberg tumours but the term has also been more broadly applied to colon, breast and pancreatic secondaries to the ovary. Combining

the data from a number of studies, in a total of 68 Krukenberg tumours, approximately 40% are metastatic from the stomach, 25% are colorectal in origin, 10% arise in the breast, and 25% arise elsewhere or do not have a primary site diagnosed. Prior to surgery many of these patients have clinical
5 and CT findings consistent with a diagnosis of ovarian cancer, and hence undergo a laparotomy. In many of these patients no evidence of another primary is found at operation, and subsequent investigations often do not reveal a primary. The pathologist may suspect that such a tumour is of non-ovarian origin based on the morphologic appearance and immunohistochemical profile,
10 but is generally not able to exclude the possibility that it could be a primary ovarian cancer, nor suggest a more likely origin. Generally these patients are given the benefit of the doubt and are treated with platinum based chemotherapy as per standard management of ovarian cancer. They usually respond poorly, and in some instances an extra-ovarian primary becomes
15 apparent at a later date.

The present invention also provides a method of using a gene expression database according to the present invention for prognosis and/or diagnosis of a patient.

20

Conventional methods for treatment of cancer rely upon clinical parameters relating to anatomical site of origin, grade and spread of disease. These observations today are essentially made through such modalities as intra-operative assessment, conventional pathology through light microscopy and a
25 suite of imaging techniques. For a proportion of tumours several molecular markers can also be used to predict the behaviour of the disease or to assess the suitability of a patient for specific treatment. One example is breast tumours that express the cell surface estrogen receptor (ESR). Such patients are known to respond to treatment with the ESR antagonist tamoxifen and it is commonly
30 used as an adjuvant therapy for low grade breast cancers. For a large proportion of tumours, however, there are currently no methods for assessing such prognostic factors. Two cancer cases that may appear identical in their pathological and clinical profiles, may respond differently to chemotherapy or

radiotherapy, they may also show a different prevalence to recur and may or may not metastasise. Underlying this phenotypic behaviour are the molecular mechanisms relating to tumour development, its cellular functioning and the relationship it has with the rest of the body.

5

Using gene expression microarray analysis the activity of thousands of genes can be used to identify expression patterns related to the phenotypic behaviour. A gene expression dataset of samples that have been clinically annotated to study specific prognostic factors, relating to treatment suitability or recurrence,
10 can be used to identify the associated molecular markers or molecular pathways. Therefore, similar to the application for identifying site of origin, where tissue differentiation markers may elude to the identity of a primary tumour, markers relating to cell survival, angiogenesis, metastasis or T-cell infiltration may be associated with tumour behaviour, patient survival or other
15 prognostic factors.

Identification of such markers or expression profiles can be translated to clinically viable tests using similar methods discussed here allowing better cancer patient management.

20

In another aspect of the present invention, there is provided a method of treatment of a patient having a tumour of unknown origin including the steps of:
identifying the tissue of origin of the tumour of unknown origin; and
treating the patient in a manner appropriate for treating a tumour
25 originating from that tissue site.

Identification of the tissue of origin permits disease-appropriate therapy to be given to a patient and thereby give the patient the best chance of receiving an effective treatment. Such treatments are known to those skilled in the art and
30 vary between different tumour origins.

Preferably, the step of identifying a tissue of origin of the tumour of unknown origin is as described herein. However, this aspect of the invention is based on

the underlying principle that an individual cell type only expresses a subset of the total number of genes present in the genome. The fraction of genes expressed reflects and determines the biological state of the cell and provides a molecular snapshot of the cellular phenotype. This is carried through to the
5 secondary or metastatic tumours and provides an identification system of their origin which allows for appropriate treatment which may not coincide with the surrounding tissue type and treatment of tumours of that tissue type.

Throughout the description and claims of this specification, the word "comprise"
10 and variations of the word, such as "comprising" and "comprises", is not intended to exclude other additives, components, integers or steps.

The discussion of documents, acts, materials, devices, articles and the like is included in this specification solely for the purpose of providing a context for the
15 present invention. It is not suggested or represented that any or all of these matters formed part of the prior art base or were common general knowledge in the field relevant to the present invention as it existed in Australia before the priority date of each claim of this application.

20 Examples of the procedures used in the present invention will now be more fully described. It should be understood, however, that the following description is illustrative only and should not be taken in any way as a restriction on the generality of the invention described above.

25 **EXAMPLES**

Example 1: Creating a Gene Expression Database.

A training dataset containing the gene expression measures of approximately 10,000 genes in a wide range of human tumour types was created. To develop the dataset, and also to ensure its usefulness for diagnosing tumour
30 type from small biopsies, a protocol incorporating an amplification step in preparation of labelled cDNA for hybridisation was used. The protocol reliably produced expression data from 3µg of starting total RNA. Amplification was an important approach to take, as the amount of tissue available is often limited to

small amounts in excess of tissue required for other diagnostic purposes. In particular, the approach allows utilising small biopsies (for example core biopsy or fine needle aspirate) of tissue collected from metastatic deposits that would otherwise not be collectable by excision biopsy.

5

a) Collection of tissue samples

All human tumour material was collected and used in accordance with the Ethical Principles as described in the Australian National Health and Medical Research Council National Statement on Ethical Conduct in Research Involving
10 Humans. Histopathology of the tumour samples was reviewed to ensure an unequivocal clinical diagnosis. Metastatic tumours arising from known primary tumour were obtained from patients with clear clinical history of metastatic disease. Pathology review of these samples unequivocally identified the primary site. Metastatic tumour arising from an unknown primary tumour was
15 submitted after substantial clinical workup. Immunohistochemical and morphological staining and review were carried out according to standard protocols.

b) Total RNA preparation and labelling

20 Tissues samples were homogenised in Trizol reagent (Invitrogen) followed by phase separation and subsequent purification of Total RNA using an RNeasy column (Qiagen) according to the manufacturers' protocols. mRNA was then amplified using standard techniques. Briefly, mRNA was reverse transcribed to cDNA using a T7 promoter tagged anchored PolyT primer. A second strand
25 was synthesized in the presence of RNaseH and Klenow. The resulting double stranded molecules were used as template in an in vitro transcription reaction using a T7 Megascript kit (Ambion), according to the manufacturer's protocol, and purified using an RNeasy column. Amplified RNA was indirectly labeled by incorporation of amino-allyl dUTP (Sigma) during reverse transcription followed
30 by coupling of cyanine-5 fluorophor (Amersham). A common reference RNA containing eleven human tumour cell line RNAs was used in all hybridisations. Reference total RNA was isolated, amplified and labeled with cyanine-3 fluorophor (Amersham) in an identical manner to the tumour samples. Samples

of labeled cDNA were cohybridised to spotted cDNA microarrays containing approximately 10,500 elements representing 9,389 unique cDNAs (UniGene build 144), washed and scanned (Scanarray 5000, Perkin Elmer) according to standard protocols. Data was extracted from scanned images using the
5 Quantarray program (GSI Luminomics).

Example 2: Profiling a tumour sample.

Samples of RNA from 121 well characterized tumour samples were analysed. To ensure the authenticity of the gene expression profiles and not to introduce
10 errors into the class prediction algorithm, the diagnosis of these samples was verified by histopathology prior to inclusion in the study. RNA from tumour samples was isolated, amplified, and labelled, and the resulting labelled cDNA was hybridised to a spotted cDNA microarray containing 9,389 unique genes (UniGene build 144). After filtering to remove unusable spots, the data were
15 normalized. Unsupervised hierarchical clustering using all genes in the filtered and normalized dataset showed the tumours grouped into their tissue of origin (Figure 2), although not perfectly. This is a not an unexpected observation and is in agreement with other studies of a similar type. A list of genes that were significantly different in expression ($p < 0.05$) between all the different tumour
20 groups was then identified using the normalization technique and informatic tools such as k-nearest neighbours and SVM. Hierarchical clustering of the samples using these genes showed significant clustering of most members of the tumour groups (Figure 3). Some tumour groups were distinct from every other tumour type (for example prostate), while others were initially more
25 difficult to separate (lung, breast, ovarian). This most likely reflects the heterogeneity of the samples, and is overcome by increasing the representation of these tumour types.

An algorithm for identifying the origin of carcinoma of unknown primary was
30 implemented, which utilises a number of informatic tools including k-nearest neighbours and a support vector machine approach. The first stage is to reduce the number of genes from the approximately 9,389 unique genes on the microarray to an optimal subset, capable of reliably describing differences

between tumour types. The optimal number and selection of genes for classification of tumours from a range of primary origins is determined by using an iterative signal to noise ratio algorithm. This method ranks genes according to the difference of their mean expression values for each class of tumour, divided by the sum of the standard deviations, ie. $(m_1 - m_2)/(s_1 + s_2)$. This effectively identifies those genes that have a consistently different expression measurement within a given class of tumours, relative to the values of that gene across all other tumour types present. A subset of such genes is shown in Figure 3. Genes are ranked according to this measurement and varying numbers of genes are tested to identify a subset with the highest predictive strength (Figure 4).

A leave-one-out (LOO) cross validation in conjunction with the k-nearest neighbors algorithm to select and test subsets of genes was then used. Briefly, this algorithm seeks to classify an unknown sample by comparing it to samples of known class by using a distance metric. The class of the closest 'k' samples is assigned to the sample being tested. LOO involves permutations of the dataset whereby each sample is held out separately and a class assigned to it by using the remaining samples. This is repeated until each sample has been left out of the training set once and has been assigned to a class. The proportion of correct classifications is used as a measure of predictor accuracy. From these predictions a confusion matrix can be constructed, as shown in Figure 5. By plotting the actual tumour classes on one axis and the predicted classes on the other, a histogram-type view of the overall success or failure of the classification approach can be achieved. This representation also allows identification of any particular classes with more incorrect predictions relative to other tumour types. The average prediction accuracy in LOO analysis in our training set is approximately 97%.

Techniques such as clustering and class-prediction algorithms are sensitive to systematic differences between samples, for example the quality of RNA, or the preparation method. To verify that amplification did not introduce errors into the dataset, the fidelity of amplification by comparison of results derived from

amplified and unamplified starting material was determined. The correlation between amplified and unamplified results was typically greater than 0.85. Between amplified samples, the correlation was greater, with a correlation coefficient of at least 0.97 (data not shown). We believe therefore that the class
5 predictions made by this algorithm are unlikely to be influenced by amplification of mRNA derived from samples.

To test the validity of the prediction algorithm we used it to identify the origin of twelve samples of metastatic tumour from a known primary. All metastases
10 were assigned to their correct class, ie known site of origin. p-values were significant in 10 cases ($P < 0.05$) and bordering on significant ($p = 0.057$, and $p = .058$) in the remaining two (see Figure 6). These specimens were not involved in any way in the construction of the prediction algorithm, and demonstrate that the prediction method is not specific (or 'over fitted') to samples contained in the
15 training set of tumours and reflects classifications based on gene expression inherent to the tumour types.

Example 3: Diagnosis of metastatic tumour in the ovary and identification of extra-ovarian origin.

20 To demonstrate the wider utility of this approach to diagnosing metastatic tumour in the ovary, we analysed three samples of tumours from the ovary which were atypical presentations suggestive of an extra-ovarian origin for the tumour. Expression data from these samples strongly suggested a colorectal origin for these tumours ($p < 0.001$ in all cases). Using only the unequivocally
25 diagnosed ovarian and colorectal tumours in the training dataset, we identified a list of 55 genes which were significantly different between the ovarian and colorectal tumours. Importantly, several genes already known to be discriminators between these tumour types were included in the list. Using just these 55 genes, the five cases described above were clearly identified as
30 colorectal in origin, and not unexpectedly, all ovarian and colorectal tumours were correctly segregated. We suggest that these genes are likely to be extremely useful as discriminators between colorectal and ovarian tumour in cases where the diagnosis is unclear or uncertain.

Example 4: Case studies and diagnosis of primary tumours**a) Colorectal primary and ovarian secondary**

A patient (P00819, Figure 7) presented with a large left ovarian mass. While
5 the clinical picture was thought to be consistent with a possible primary ovarian
cancer, this patient had presented with a Duke's C colon carcinoma one year
previously. She underwent surgery and the histology was initially reported as a
moderately differentiated mucinous adenocarcinoma with light microscopic
appearances favouring a primary left ovarian cancer with omental involvement.
10 Immunohistochemical analysis revealed a phenotype more consistent with a
colonic metastasis, as the tumour was found to be CK 7 negative and CK 20
positive. This illustrates the scenario that we expect to frequently encounter
where the clinical picture, diagnostic pathology and PET imaging suggest a
primary tumour location, but without a high degree of certainty and with some
15 conflicting data regarding the origin of the tumour. In this case, molecular
profiling was of immediate applicability, and supported the immunophenotyping
of this tumour as colorectal.

b) Colorectal primary and pelvic and peritoneal secondary

20 The patient (P00644, Figure 7) presented with a pelvic tumour mass and
widespread peritoneal metastases. Surgical notes from the time of operation
indicated it was unclear whether the patient had a primary ovarian or a primary
colorectal cancer. Histology of the tumour was reported as a moderately
differentiated endometrioid adenocarcinoma with some focal mucinous
25 differentiation, with the light microscopic appearances favouring ovary as the
primary. Immunohistochemical staining with CK7 and CK20 monoclonal
antibodies showed tumour cells variably co-expressing the two markers, which
was thought to support an ovarian origin, although without a high degree of
certainty. In this case, our molecular profiling suggested that the likely true
30 origin of the tumour was colorectal.

c) Colorectal primary and ovarian secondary

Further samples of tumours isolated from the ovary, where an extra-ovarian origin for the tumour was likely were examined. The first (P00482, Figure 7) was a sample collected from the ovary of a woman with abdominal metastases at the time of left hemicolectomy, total abdominal hysterectomy and bilateral salpingo-oophrectomy. Molecular profiling identified a colorectal origin for the tumour, which was confirmed by histological analysis of sections of the colon, which showed that the patient had a Duke's stage D moderately differentiated adenocarcinoma of the sigmoid colon. The second patient (P00493, Figure 7) presented with tumour present in both ovaries, and omentum. She had previously been treated for carcinoma of the sigmoid colon, and clinicians queried whether the tumour was a recurrence from the colorectal tumour, or an ovarian primary. In this case, microarray analysis indicated colorectal as the likely source of the tumour, which was confirmed by immunohistochemical staining which showed negative staining for cytokeratin 7, and positive staining for cytokeratin 20. The third patient (P00206, Figure 7) was never diagnosed as a colorectal tumour metastatic to the ovary, although at the time of surgery, the pathologist noted, "although the (histologic) appearances would be consistent with mucinous ovarian carcinoma the appearances nevertheless raise the possibility of this representing spread from a colorectal primary tumour."

20

Example 5: Various uses of the microassay.

We expect that this test will be useful in a number of clinical situations. The first involves a patient presenting with no previous history of cancer, with extensive undifferentiated carcinoma. This is the classical presentation of carcinoma of unknown primary. In one such case (P00459), we analysed a sample of a carcinoma taken from a forty year-old non-smoker who presented with cough and dyspnoea. The patient was subsequently found to have multiple lung, supraclavicular, mediastinal and liver metastases. Histology review of the metastatic tumour described as an undifferentiated carcinoma. There was a larger lesion in the right lung on CT that may have been consistent with a primary. A PET scan did not reveal a definite primary, although a questionable abnormality in the lower oesophagus was noted. Subsequent gastroscopy was normal. Although the clinical picture was consistent with a diagnosis of a non-

30

small cell lung cancer, there remained considerable uncertainty about the primary origin of this cancer in a young non-smoker. Expression profiling of this sample, and subsequent comparison with the training dataset determined that this sample had an expression profile most consistent with the tumour being
5 lung in origin, with a significant p-value of 0.027. This case illustrates the scenario where the clinical picture, diagnostic pathology and imaging suggested a primary tumour location, but with some remaining doubt. Array analysis subsequently confirmed the clinical observations.

10 The second scenario we expect to encounter frequently is the unusual presentation of a common tumour, when that patient has a clinical history of a previous cancer. One of the patients in our study (P00563), a thirty-one year old woman with a past history of a stage I high-grade mucinous borderline ovarian tumour six years previously, presented with a twelve month history of
15 left pelvic pain and was found to have a sclerotic abnormality involving the left ilium and left upper femur. Bone scan revealed multiple bone metastases. Biopsy from the left ilium revealed adenocarcinoma. The patient underwent CT scan of the chest/abdomen/pelvis, a PET scan, a thallium scan and a mammogram without any evidence of a primary being found. Pathology review
20 suggested that the histology was consistent with a previous ovarian malignancy, but could not exclude a carcinoma arising in the breast, lung or gastrointestinal tract. The presentation with bone metastases was thought to be most unusual for recurrent ovarian cancer and the treating clinician thought that it was more likely that the cancer had arisen from another site. The patient was treated as
25 an unknown primary with a combination of epirubicin, cisplatin and 5-fluorouracil. Our array analysis confirmed the possible diagnosis of a relapse from the ovarian primary, and it is possible that information such as this may have altered the management of this patient.

30 The third scenario involves a patient with a clear history of malignancy, but with metastatic tumour where it is unclear whether the metastatic tumour has arisen from the first, or a new, primary tumour. In some cases, we expect that array analysis would be able to confirm the identification of a relapsed primary

tumour, and in others to suggest a new primary site. Both of these scenarios were encountered during this work. The first was a patient (P00563) diagnosed in February 1994 with Stage IIC endometrioid carcinoma of the ovary. CA125 was elevated at 327 pre-operatively and was still elevated post-operatively at 5 80. She underwent a total abdominal hysterectomy with bilateral salpingo-oophorectomy and omentectomy. She was then treated with six cycles of carboplatin and cyclophosphamide. She remained well until May 1998 when she developed back pain and was found to have sclerotic bone metastases (investigations included plain x-ray, CT and bone scan). Mammogram at this 10 time was normal. The CT scan did not reveal any other evidence of metastatic disease. T9 metastasis was biopsied and revealed a poorly differentiated adenocarcinoma, which was oestrogen and progesterone receptor negative. There was no clinical evidence of another primary. Although the development of sclerotic bone metastases was thought to be an unusual pattern of relapse 15 for ovarian cancer, the decision was made to treat her as ovarian cancer. In addition to the CT she also had a PET scan, which was unhelpful. Following palliative radiotherapy to thoracic spine, she went on to receive six cycles of carboplatin and taxol. The response was difficult to assess though there was some slight improvement on bone scan. The CA125 tumour marker was not 20 elevated and never rose subsequently. By October 1999 there was a definite mass in the left neck and initial attempt to biopsy this mass in January 2000 did not reveal any malignant cells. In April 2000, due to progressive growth and symptoms from the neck mass the patient received palliative radiotherapy, and commencing in June 2000 received four cycles of carboplatin with the best 25 response of stable disease. In November 2000, there was an impression of a mass in the outer left quadrant of the left breast with some suspicious changes of malignancy on mammogram in the same area. However, biopsy of the breast mass was negative. The patient also had a repeat biopsy of the neck mass, which revealed an undifferentiated carcinoma. At this time she was also 30 thrombocytopenic and had developed liver metastases. Bone marrow examination revealed a similar undifferentiated carcinoma. She was commenced on capecitabine but tolerated this poorly and this treatment was ceased. She subsequently went on to receive weekly taxotere with

improvement in her thrombocytopaenia but progressive liver metastases. She was subsequently treated palliatively, and died in May 2001. At the time she was initially found to have bone metastases she was regarded as an atypical relapse of ovarian cancer but there was always concern that she may have had another primary, in particular, a breast primary in view of the sclerotic bone metastases. By November 2000, there was a strong clinical suspicion that she may have had breast cancer, and this influenced the decision to use capecitabine and docetaxel chemotherapy. Although the biopsy of the breast mass was negative this was a poor sample and may have been a false negative. The clinical and mammographic appearances of the breast lesion were consistent with a breast primary, as was the pattern of metastases with liver, bone marrow, left supraclavicular and sclerotic bone metastases. Analysis of this sample by microarray, subsequent to the patient's death confirmed the suspicion that the metastatic cancer was not a relapse of the patient's initial ovarian cancer, but a new breast primary.

The alternative scenario, where a relapse was suggested, involved a seventy year old man (P01242) who was diagnosed with prostate cancer ten years previously and treated by transurethral resection of the prostate, radiotherapy and had remained on Zoladex and flutamide. He presented with a painful lesion on his left ear and a lump in the left upper neck. Biopsy of the ear lesion revealed no evidence of malignancy, but initial core biopsy of the hard 1.5 cm left upper neck mass was reported as poorly differentiated metastatic carcinoma, and immunohistochemistry for PSA was negative. Serum PSA was also normal. He was referred to another hospital for further investigation. A repeat biopsy was performed and our analysis of a sample by molecular profiling identified prostate as the likely source of the tumour. This biopsy was initially reported as metastatic adenocarcinoma with focal neuroendocrine differentiation, and the pathologist recommended that a lung primary should be excluded. Repeat immunohistochemical staining for PSA on this biopsy was requested after the array result was already known, and this was positive consistent with metastatic prostate cancer.

The data presented shows that the use of expression profiling is able to contribute to the management of cancer patients. This work demonstrates that whilst expression changes may occur in some genes as a result of tumour development, or admixing of cells with other cell types such as stroma or vascular elements, the mass effect of measuring the expression patterns of thousands of genes means that distinctive patterns of tumour types are identifiable. Further, expression profiles are shown to be sufficient to classify tumour samples according to tissue of origin. It has been demonstrated that, not only can tumours be partitioned with respect to tissue of origin using microarray analysis, but additionally the expression patterns can be used to positively identify samples which were previously unknown.

Example 6: Use of RT-PCR.

a) Extraction of RNA from Paraffin Embedded Formalin Fixed Tissue (FFPET)
Extraction of RNA was performed using a modification to the protocol described by Specht *et al* (2001, *Am J Pathol* 158(2): 419-29). Briefly, paraffin was removed from microtome sectioned material by incubating in Xylene, repeating the procedure twice and then sequentially washing with 100%, 90% and 70% ethanol. Samples were then dried before the addition of Proteinase K digestion buffer (10mM TrisHCl (pH 8.0), 0.1mM EDTA (pH8.0), 2% SDS), and 100 mg of Proteinase K, followed by incubation at 60°C for 16 hours or overnight. Following the initial incubation period an additional 100 mg of proteinase K was added and samples were digested for a further 3-4 hours at 60°C. RNA was purified from the tissue lysate by column chromatography (Rneasy, Qiagen) using a modification to the manufacturers protocol. This involved sequentially adding 440 µL of 100% ethanol and 660 µL of buffer RLT buffer to the tissue lysate. The sample was then briefly mixed before passing it through the column by centrifugation (RNeasy mini). Subsequent washes were applied as per described by the manufacturer followed by elution in an appropriate volume of RNase free deionised water.

b) Quantitative real time PCR

Total RNA was reverse transcribed by priming with random hexamers. Success of the reverse transcription and relative quantification of the cDNA was interpreted using 5 endogenous control genes, analysed by real time PCR using SYBR green chemistry (ABI Prism 7000). The endogenous control genes
5 CTCF, CAPZB, TXNL, SF3B1, RABGGTB and PGK were chosen from microarray experiments based on the criteria of low variability across multiple cancer classes and a minimum expression level in excess of three times that of background. All primer pairs were designed across exon boundaries to prevent amplification of genomic DNA. An average of the Ct values for the endogenous
10 controls was used to assess the quantity of cDNA present. A maximum average Ct threshold was set to exclude samples not suitable for further analysis on micro fluidics card.

c) Micro Fluidics Card

15 A set of 89 genes was chosen by signal to noise gene selection using a 6 class training set of breast, colorectal, ovarian, gastric, pancreas and a combined class (others) representing other sites of origin (ie lung, melanoma, prostate, renal, mesothelioma, testicular, SCC). The genes represent the top ranked 12 to 17 markers for each respective class by signal to noise gene selection. All
20 genes were chosen from Applied Biosystems Assay on Demand (AoD) pre-validated primer probe sets. If a gene marker selected by the signal to noise metric was not available from the AoD set then the next highest ranking gene was selected. Additionally, seven endogenous controls were added to the assay set including the 5 genes previously described for cDNA quality control
25 and mandatory controls 18s rRNA and GAPDH. Custom microfluidics cards were designed in a configuration allowing the processing of 4 samples and 96 assays on a single card.

A master mix of reagent was prepared from TaqMan® Universal PCR Master
30 Mix and sample cDNA template. The volumetric amount of template used was proportional to that used for quality control with no attempt to standardise the absolute amount of template added between samples. Reactions were run according to the manufacturers protocol with data collection based on absolute

Ct values. Normalisation of RT-PCR assays was conducted using an average Ct value for all endogenous controls excluding GAPDH. Samples were then converted to a fold ratio relative to endogenous controls described using standard delta Ct formula.

5

i.e. $X = 2^{\Delta Ct}$ where $\Delta Ct = (Ct_{\text{target}} - Ct_{\text{average endogenous controls}})$

Example 7: Generation of gene expression database – validation of RT-PCR results.

- 10 A cohort of 42 samples spanning five anatomical sites of origin (breast, colorectal, gastric, pancreas, ovarian) was profiled using RT-PCR by custom micro fluidics cards. All reactions were performed using cDNA generated from RNA extracted from fresh frozen tissue. These samples had been previously analysed using cDNA microarrays. A comparison of median normalised data by
- 15 heat map alignment shows the consistency between the two platforms (Figure 9).

The chemistry used for RT-PCR analysis allows the utilisation of nucleic acids that may be partially degraded or fragmented, as opposed to microarray

20 analysis where high quality intact mRNA is required. Formalin fixation of tissue is routinely used in conventional pathology to conserve tissue architecture and preserve protein complexes that may be targeted by immunohistochemical detection as cancer specific markers. The cross-linking events that allow this preservation, however, are detrimental to RNA and DNA integrity. Nucleic acids

25 extracted from such material are therefore composed of short fragments, typically of around 300 bp in length. RT-PCR requires the amplification of only short lengths of DNA. Amplicon lengths generated from AoD primer sets are approximately 60 bp in length.

- 30 Applicants have used RNA extracts from FFPET for expression profiling using RT-PCR using the micro fluidics format. A total of 13 samples from 5 sites of origin were processed providing high quality data. Clustering of samples processed from both fresh frozen tissue and FFPET show that samples can

accurately be grouped into respective tumour classes regardless of the tissue processing method used prior to RNA extraction (Figure 10).

5 Similar to microarray data, data generated from RT-PCR can be used for machine learning and creating class predictor models. All RT-PCR data was used for generating an SVM predictor model of 5 classes (breast, gastric, ovarian, colorectal and pancreas) using the method of ranking. Using 5 RankLevels applicants achieved a LOO cross validation accuracy of 100%. The versatility of a rank method for cross platform meta-analysis was also applied to both microarray and RT-PCR datasets. Training solely using data generated by cDNA microarray SVM models were generated that can be tested upon similar samples profiled using RT-PCR. Using this cross platform meta-analysis a high prediction accuracy of 93% was obtained in the independent test.

15

Example 8: Testing Strengths of Predictions

The strength of the prediction capability for a carcinoma unknown primary (CUP) was tested. This test is indicative of whether a prediction of the tissue of origin for a carcinoma of unknown primary is correct. When a class or histological subclass is left out of a training set used to establish the gene database the prediction accuracy of the test is compromised. This demonstrates the importance of having all classes or subclasses present when establishing a training set.

25

The present example tests the veracity of the prediction strength algorithm, and associates a confidence with the prediction.

30 Figure 12 shows that data set size has an impact on the confidence of the prediction. By changing the number of samples in the dataset available for comparison, the degree of confidence is affected. Lowering the number or leaving out data sets reduces the confidence level.

Finally it is to be understood that various other modifications and/or alterations may be made without departing from the spirit of the present invention as outlined herein.

CLAIMS

1. A method of profiling a biological sample, said method including:
 - obtaining a gene expression profile from the biological sample;
 - 5 obtaining a gene expression database from one or more biological samples;
 - identifying different patterns of gene expression between the biological samples;
 - identifying genes that comprise the different patterns of gene expression; and
 - 10 correlating the genes that comprise the different patterns of gene expression of the gene expression profile of the biological sample and the gene expression database to provide a profile of the biological sample.
- 15 2. A method according to claim 1 wherein the patterns of gene expression are normalized by using an iterative signal to noise ratio algorithm wherein:

$$(m_1 - m_2) / (s_1 + s_2)$$

and where m = mean expression value; and
 s = standard deviation.
- 20 3. A method according to of claim 1 or 2 wherein the gene expression is processed by a formula for normalization according to the formula:

$$X = 2^{\Delta Ct} \text{ where } \Delta Ct = (Ct_{\text{target}} - Ct_{\text{average endogenous controls}})$$
- 25 4. A method according to any one of claims 1 to 3 wherein the different patterns of gene expression are identified by an analysis which employs an algorithm utilising a k-nearest neighbours and/or support vector machine (SMV) approach.
- 30 5. A method according to any one of claims 1 to 4 wherein the patterns of gene expression are further clarified using leave-one-out (LOO) cross validation in conjunction with a k-nearest neighbour algorithm and/or SVM.

6. A method according to any one of claims 1 to 5 wherein the gene expression profiles and the gene expression databases include gene expression data that is processed by ranking genes according to their expression levels within a sample and allocating a rank to the gene such that
5 the rank of the gene identifies different patterns of gene expression between the biological samples.

7. A method according to claim 6 wherein the rank is allocated a rank level using the following formula:

10 **RankLevel** = *ceil* (number of rank levels x rank of the gene/number of genes assayed) = *ceil* (x) wherein *ceil* (x) = smallest integer \geq x

8. A method according to any one of claims 1 to 7 wherein the gene expression profile includes gene expression data obtained by analysis of gene
15 expression from the biological sample by at least one method selected from the group including RNA expression analysis, DNA expression analysis and transcription rate analysis.

9. A method according to any one of claims 1 to 8 wherein the analysis of
20 gene expression is obtained by RNA expression analysis.

10. A method according to any one of claims 1 to 9 wherein the analysis of gene expression is obtained by a hybridisation based method or a PCR based method.
25

11. A method according to any one of claims 1 to 9 wherein the analysis of gene expression is obtained by microarray analysis.

12. A method according to any one of claims 1 to 9 wherein the analysis of
30 gene expression is obtained by quantitative RT-PCR.

13. A method according to any one of claims 1 to 9 wherein the analysis of gene expression is obtained by microarray and quantitative RT-PCR.

14. A method according to any one of claims 1 to 13 wherein the biological sample is selected from the group including normal tissue, a pre-cancerous, cancerous or tumour tissue, a primary tumour sample, cells from pleural effusions, or metastatic samples.
15. A method according to any one of claims 1 to 14 wherein the tumour tissue is from a tumour of unknown origin.
16. A method according to any one of claims 1 to 15 wherein the gene expression database is generated from a plurality of gene expression profiles of biological samples and/or samples of tumour types.
17. A method according to claim 16 wherein the tumour types are selected from the group including gastric, colorectal, pancreatic, breast and ovarian.
18. A method according to claim 17 wherein the tumour samples or tumour types are selected from Table 1.
19. A method according to any one of claims 1 to 18 wherein the gene expression profile or the gene expression database includes an analysis of a gene selected from Table 2.
20. A method of processing gene expression data obtained from more than one gene expression analysis of a biological sample, said method including ranking genes according to their expression ratios within a sample processed by the gene expression analysis and allocating a rank to the gene such that the rank of the gene identifies a different pattern of gene expression between the biological samples.
21. A method according to claim 20 wherein the rank is allocated a rank level using the following formula:

RankLevel = *ceil* (number of rank levels x rank of the gene/number of genes assayed) = *ceil* (x) wherein *ceil* (x) = smallest integer \geq x

22. A method according to claim 20 or 21 wherein the gene expression data
5 is normalised using an iterative signal to noise ratio algorithm wherein:

$$(m_1 - m_2) / (s_1 + s_2)$$

and where m = mean expression value; and
s = standard deviation.

- 10 23. A method according to any one of claims 20 to 22 wherein the gene expression data is processed by a formula for normalization according to the formula:

$$X = 2^{\Delta Ct} \text{ wherein } \Delta Ct = (Ct_{\text{target}} - Ct_{\text{average endogenous controls}})$$

- 15 24. A method according to any one of claims 20 to 23 wherein the gene expression data has been analysed using an algorithm utilising a k-nearest neighbours and/or a support vector machine (SMV) approach.

25. A method according to any one of claims 20 to 24 wherein the gene
20 expression data is further clarified using leave-one-out (LOO) cross validation in conjunction with a k-nearest neighbour algorithm and/or SVM.

26. A method according to any one of claims 20 to 25 wherein the gene
expression data is obtained by analysis of gene expression from the sample by
25 at least one method selected from the group including RNA expression analysis, DNA expression analysis and transcription rate analysis.

27. A method according to any one of claims 20 to 26 wherein the analysis of
gene expression is obtained by RNA expression analysis.

30

28. A method according to any one of claims 20 to 27 wherein the analysis of
gene expression is obtained by a hybridisation based method and/or a PCR
based method.

29. A method according to any one of claims 20 to 27 wherein the analysis of gene expression is obtained by microarray analysis.

5 30. A method according to any one of claims 20 to 27 wherein the analysis of gene expression is obtained by quantitative PCR.

31. A method according to any one of claims 20 to 27 wherein the analysis of gene expression is obtained by microanalysis and quantitative PCR.

10

32. A method according to any one of claims 20 to 31 wherein the biological sample is selected from the group including normal tissue, a pre-cancerous, cancerous or tumour tissue, primary tumour and metastatic tumour, or cells from a pleural effusion.

15

33. A method according to claim 32 wherein the tumour tissue is from a tumour of unknown origin.

20

34. A method according to claim 33 wherein the tumour tissue is derived from a tumour type selected from the group including gastric, colorectal, pancreatic, breast and ovarian.

35. A method according to claim 34 wherein the tumour tissue is selected from samples of Table 1.

25

36. A method according to any one of claims 20 to 35 wherein the genes are selected from Table 2.

30

37. A gene expression database including gene expression profiles from biological samples, said gene expression profiles including gene expression data obtained by analysis of different patterns of gene expression from the biological sample, said data obtained by at least one method selected from the

group including RNA expression analysis, DNA expression analysis and transcription rate analysis.

38. A gene expression database according to claim 37 wherein the analysis
5 of the gene expression is normalised by using an iterative signal to noise ratio algorithm wherein:

$$(m_1 - m_2) / (s_1 + s_2)$$

and where m = mean expression value; and
 s = standard deviation.

10

39. A method according to claim 37 or 38 wherein the gene expression is processed by a formula for normalization according to the formula:

$$X = 2^{\Delta Ct} \text{ where } \Delta Ct = (Ct_{\text{target}} - Ct_{\text{average endogenous controls}})$$

- 15 40. A gene expression database according to any one of claims 37 to 39 wherein different patterns of gene expression are identified by analysis which employs an algorithm utilising a k-nearest neighbours and support vector machine (SMV) approach.

- 20 41. A gene expression database according to any one of claims 37 to 40 wherein the different patterns of gene expression are further clarified using leave-one-out (LOO) cross validation in conjunction with a k-nearest neighbour algorithm.

- 25 42. A gene expression database according to any one of claims 37 to 41 wherein the gene expression data of the sample obtained by analysis of different patterns of gene expression is processed by ranking genes according to an expression ratio of the gene within the sample, and allocating a rank to the gene such that the rank of the gene identifies a different pattern of gene
30 expression between the biological samples.

43. A gene expression database according to claim 42 wherein the rank is allocated using a formula:

RankLevel = *ceil* (number of rank levels x rank of gene/number of genes assayed) = *ceil* (x) wherein *ceil* (x) = smallest integer \geq x

44. A gene expression database according to any one of claims 37 to 43
5 wherein the analysis of gene expression is obtained by RNA expression analysis.
45. A gene expression database according to any one of claims 37 to 44
10 wherein the analysis of gene expression is obtained by a hybridisation based method or a PCR based method.
46. A gene expression database according to any one of claims 37 to 45
wherein the analysis of gene expression is obtained by microarray analysis.
- 15 47. A gene expression database according to any one of claims 37 to 45 wherein the analysis of gene expression is obtained by quantitative PCR.
48. A gene expression database according to any one of claims 37 to 45
20 wherein the analysis of gene expression is obtained by microarray and quantitative PCR.
49. A gene expression database according to any one of claims 37 to 48
wherein the biological sample is selected from the group including normal tissue, a pre-cancerous, cancerous or tumour tissue, primary tumour or
25 malignant tumour or cells from a pleural effusion.
50. A gene expression database according to any one of claims 37 to 49
wherein the tumour tissue is from a tumour of unknown origin.
- 30 51. A gene expression database according to any one of claims 37 to 50 wherein the gene expression database is generated from a plurality of gene expression profiles.

52. A gene expression database according to any one of claims 37 to 51 wherein the tumour tissue is derived from a tumour type selected from the group including gastric, colorectal, pancreatic, breast and ovarian.

5 53. A gene expression database according to any one of claims 37 to 52 wherein the tumour tissue is selected from samples of Table 1.

54. A gene expression database according to any one of claims 37 to 53 database which includes an analysis of a gene selected from Table 2.

10

55. A method of evaluating an origin of a tumour sample, said method including:

obtaining a gene expression profile of the tumour sample;

15 comparing the gene expression profile of the tumour sample to a gene expression database said database including gene expression profiles from known tumour samples, said gene expression profiles including gene expression data obtained by analysis of gene expression from the known tumour samples by at least one method selected from the group including RNA expression analysis, DNA expression analysis and transcription rate analysis;
20 and

identifying the origin of the tumour sample when a gene expression profile from the gene expression database correlates with the gene expression profile of the tumour sample.

25 56. A method of evaluating an origin of a tumour sample, said method including:

obtaining a gene expression profile of the tumour sample according to any one of claims 1 to 19;

30 comparing the profile of the tumour sample to a gene expression database according to any one of claims 37 to 54; and

identifying an origin of the tumour sample when a gene expression profile from the gene expression database correlates with the gene expression profile of the tumour sample.

57. A method according to claim 55 or 56 wherein the origin of the tumour sample is identified when a pattern for gene expression from the gene expression database correlates to a pattern from the gene expression profile of the tumour sample.

58. A method according to claim 55 or 56 wherein the origin of the tumour sample is identified when a rank level for gene expression from the gene expression database correlates to a rank level from the gene expression profile of the tumour sample.

59. A method according to claim 58 wherein more than one rank level for gene expression from the gene expression database correlates to more than one rank level from the gene expression profile of the tumour sample.

60. A method according to any one of claims 55 to 59 wherein the tumour sample is selected from the group including a pre-cancerous, cancerous or tumour tissue, primary tumour and malignant tumour.

61. A method according to any one of claims 55 to 60 wherein the tumour sample is from a tumour of unknown origin.

62. A method of treating a patient having a tumour of unknown origin, said method including:

identifying the tissue of origin of the tumour of unknown origin; and
treating the patient in a manner appropriate for treating a tumour originating from that tissue.

63. A method of treating a patient having a tumour of unknown origin, said method including:

identifying the origin of the tumour of unknown origin according to any one of claims 55 to 61; and

treating the patient in a manner appropriate for treating a tumour originating from that tissue.

64. A microarray having a plurality of loci and having an oligopeptide affixed
5 to the loci, said oligopeptide capable of binding to a gene selected from Table 2.
65. A microarray when used for profiling a tumour according to claim 1.
66. A microarray when used for obtaining a gene expression database
10 according to any one of claims 37 to 54.

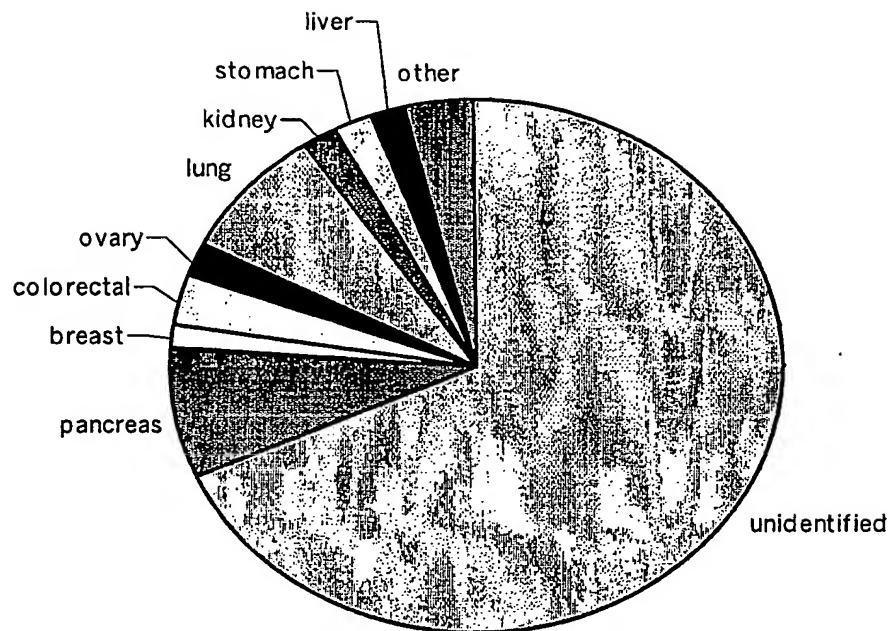
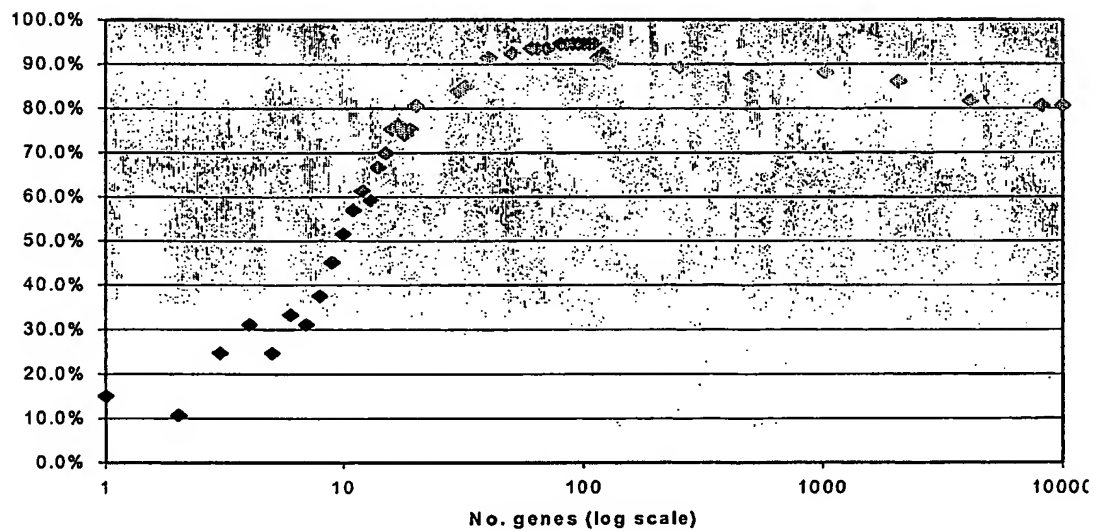
Figure 1**Figure 4**

Figure 2

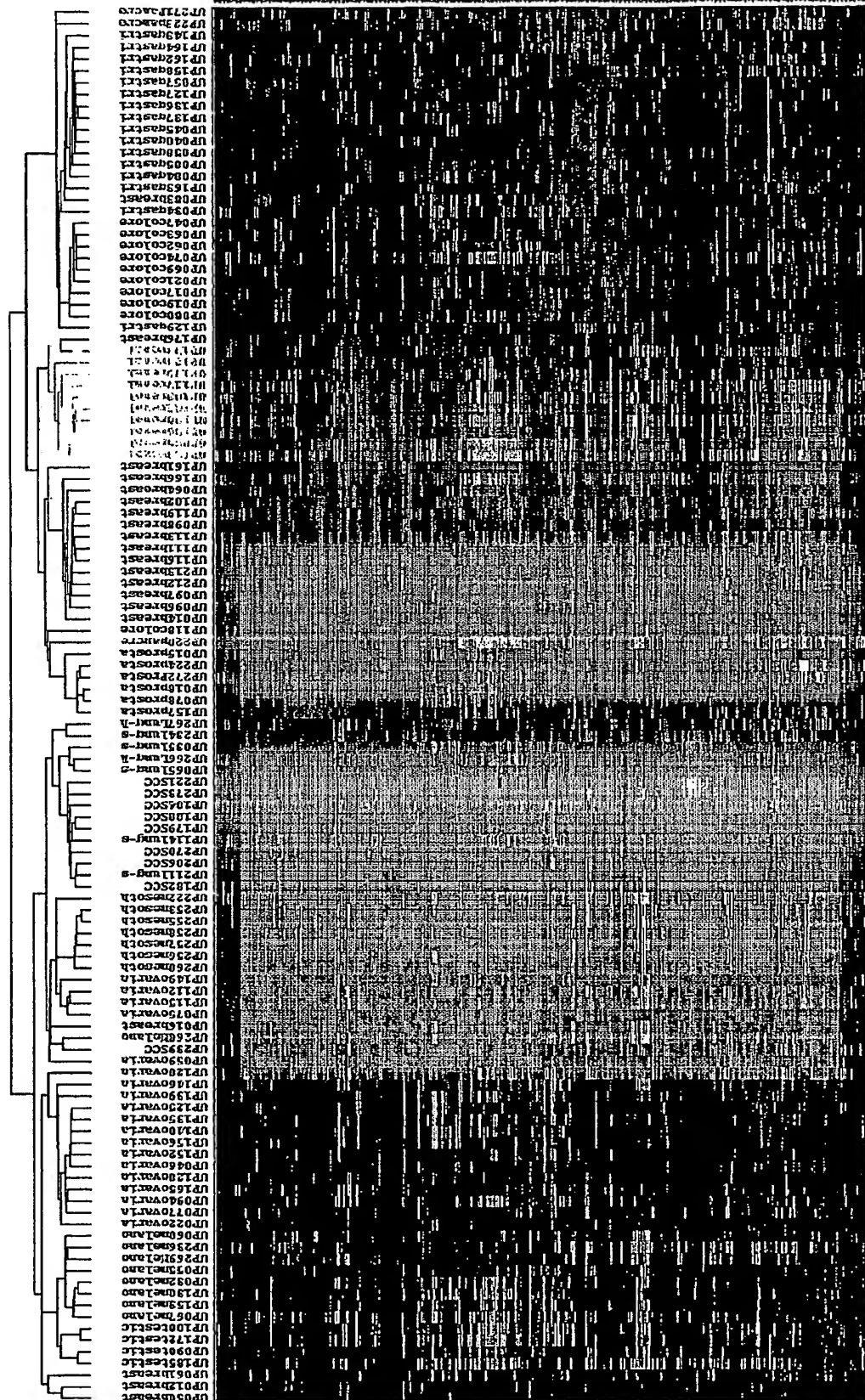
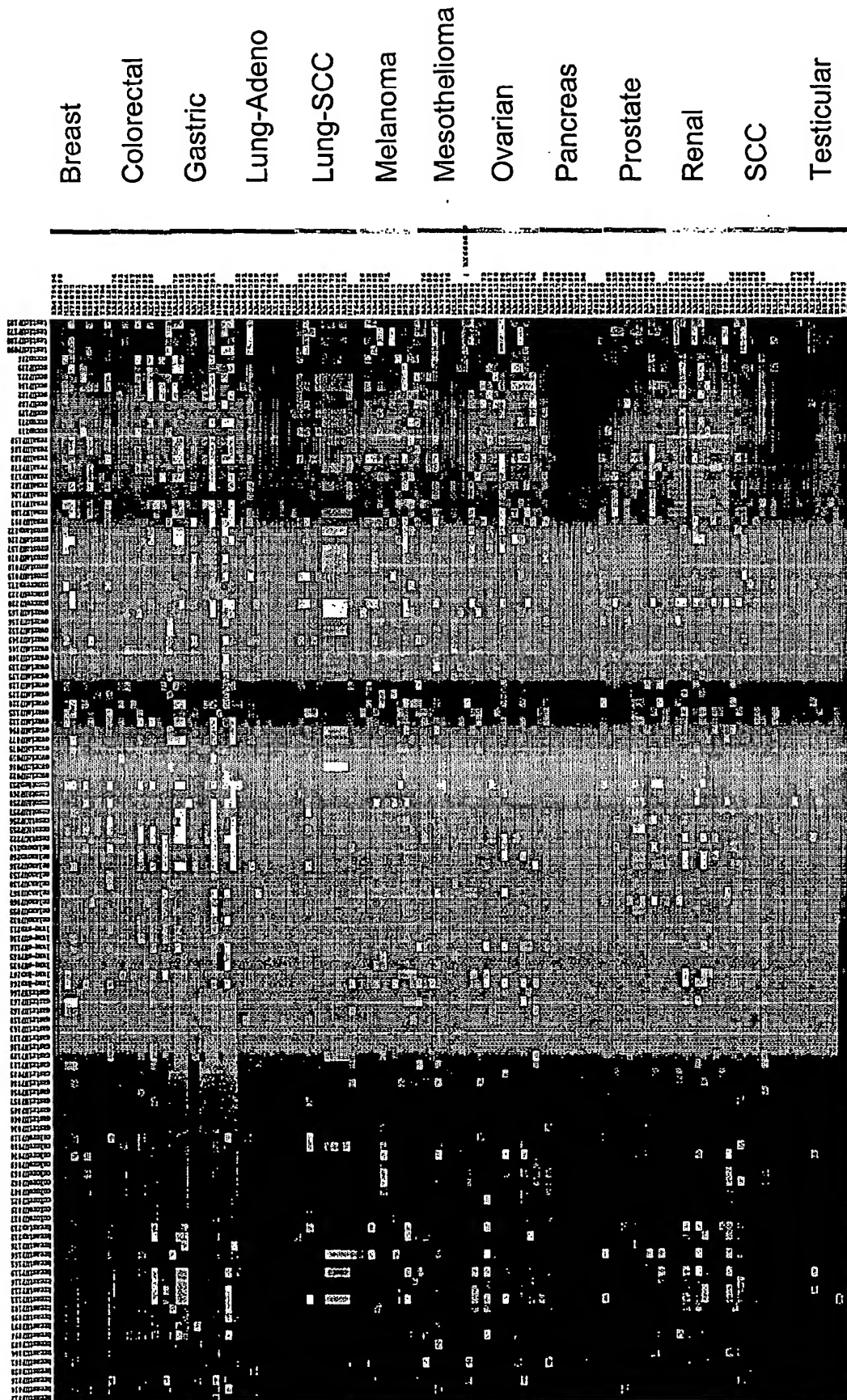


Figure 3



Breast
Colorectal
Gastric
Lung-Adeno
Lung-SCC
Melanoma
Mesothelioma
Ovarian
Pancreas
Prostate
Renal
SCC
Testicular

Figure 5

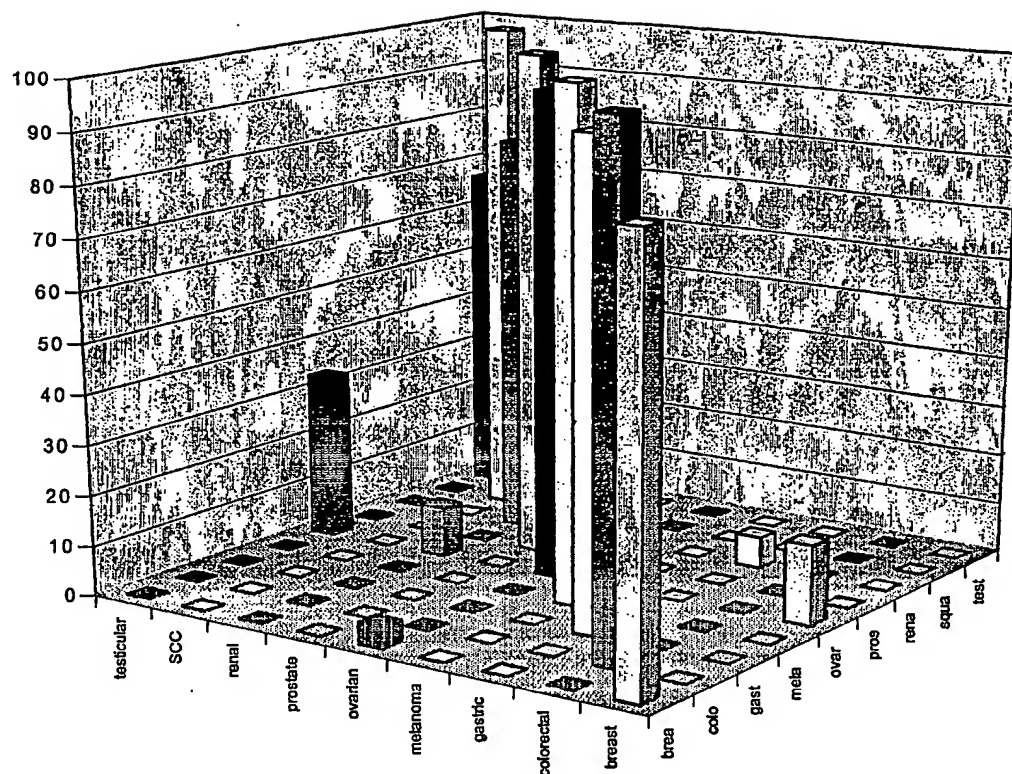


Figure 6

UP no	Type	Prediction	P-value	Comment
UP020	metastasis	colorectal	0.03734007	metastatic colorectal adenocarcinoma, liver biopsy
UP024	metastasis	colorectal	0.057077657	metastatic colorectal tumour, liver biopsy.
UP025	metastasis	melanoma	0.014119678	metastatic malignant melanoma, axillary LN biopsy
UP066	metastasis	colorectal	0.01855055	metastatic colorectal tumour, lung biopsy
sUP071	metastasis	renal	2.8264578E-4	metastatic renal cell tumour, lung biopsy
UP072	metastasis	colorectal	0.03508438	metastatic colorectal tumour, lung biopsy
UP106	metastasis	ovarian	0.024418509	metastatic papillary ovarian adenocarcinoma, omentum biopsy
UP126	metastasis	colorectal	0.0010548463	metastatic colorectal tumour, lung biopsy
UP132	metastasis	ovarian	0.042258687	metastatic serous papillary ovarian tumour, unknown biopsy
UP150	metastasis	colorectal	0.01600025	metastatic colorectal tumour, liver biopsy.
UP169	metastasis	breast	0.04593459	metastatic breast ductal carcinoma ER+, Pr+, Her2+, left ovary biopsy
UP171	metastasis	gastric	0.058362514	metastatic poorly differentiated signet ring gastric carcinoma, LN biopsy

Figure 7

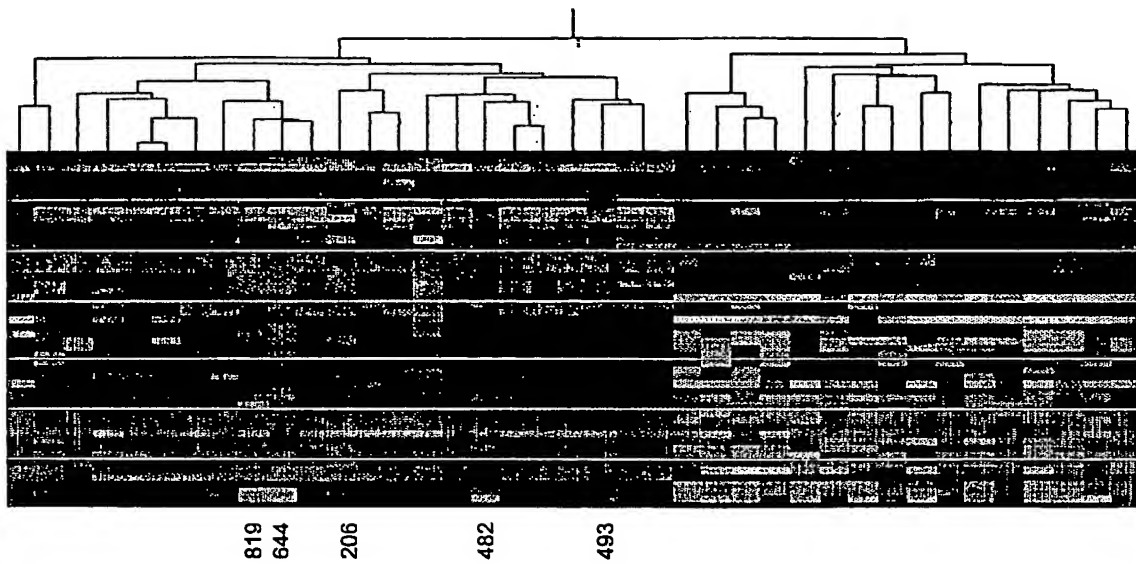


Figure 8

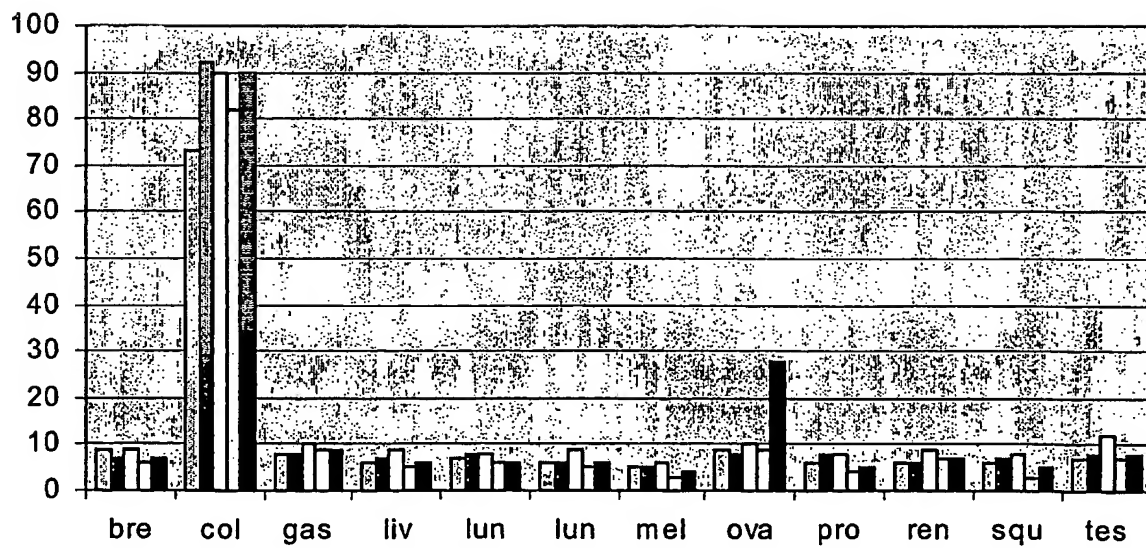
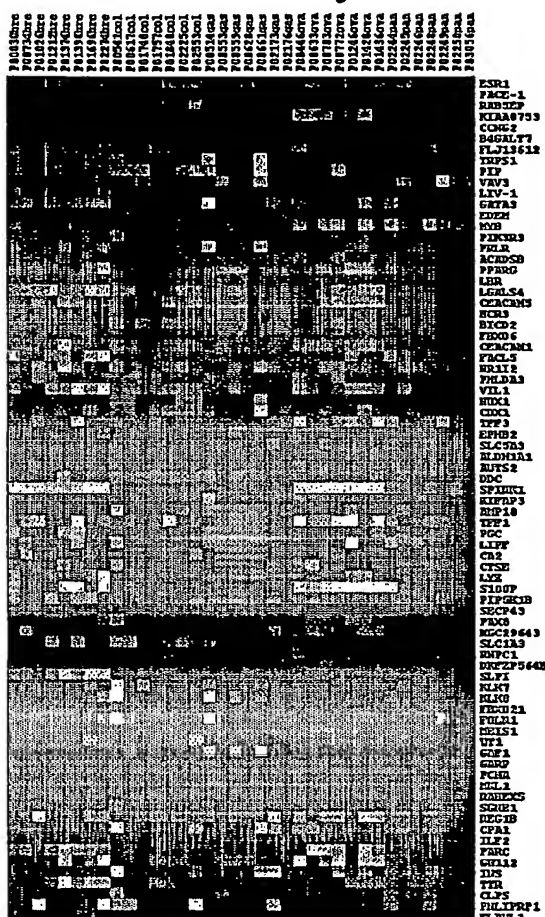


Figure 9

cDNA Microarray



RT-PCR

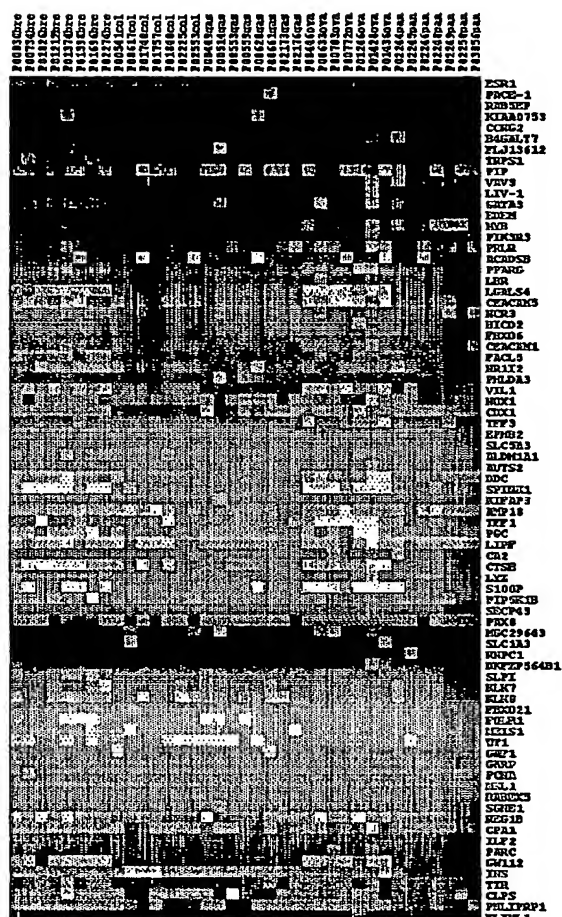
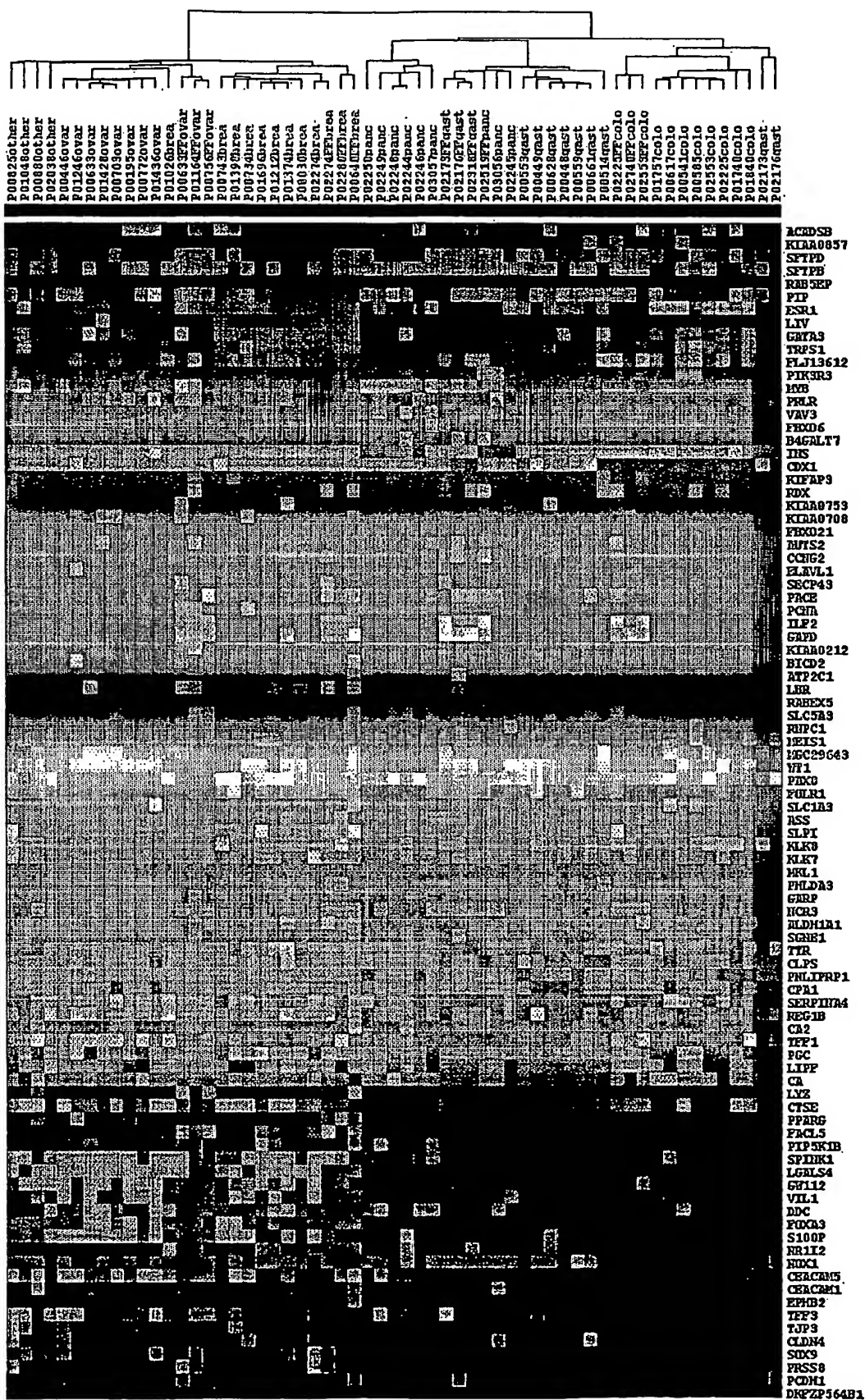


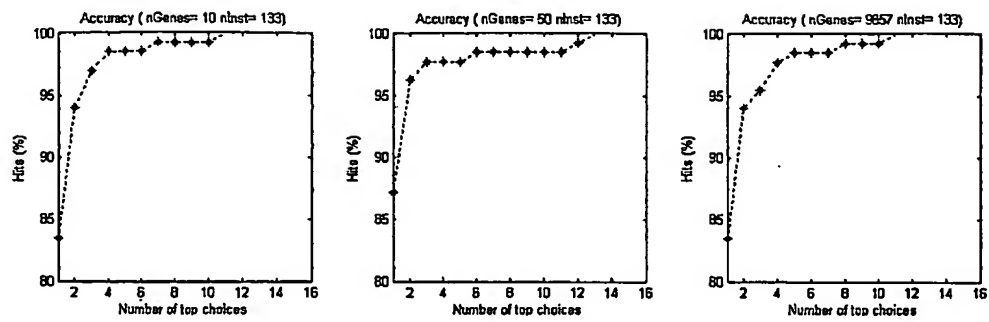
Figure 10



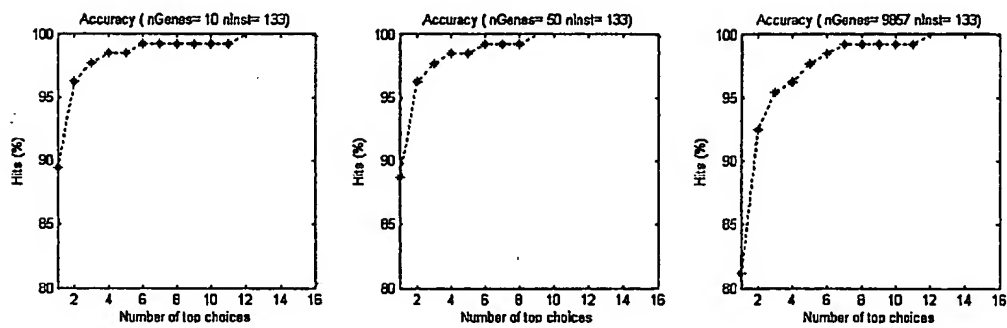
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Figure 11

A. Full precision



B. 3-RankLevels



C. 5-RankLevels

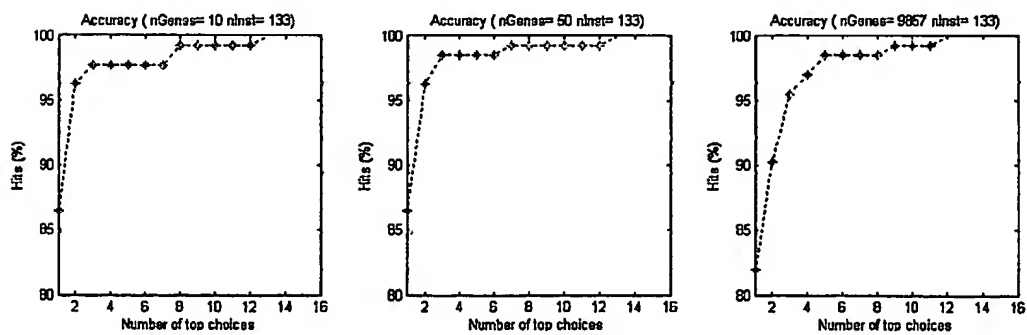


Figure 12

